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(54) Title: METHODS OF DIAGNOSIS OF PROSTATE CANCER, COMPOSITIONS AND METHODS OF SCREENING FOR MODULATORS OF PROSTATE CANCER

(57) Abstract: Described herein are genes whose expression are up-regulated or down-regulated in prostate cancer. Also described are such genes whose expression is further up-regulated or down-regulated in drug-resistant prostate cancer cells. Related methods and compositions that can be used for diagnosis and treatment of prostate cancer are disclosed. Also described herein are methods that can be used to identify modulators of prostate cancer.

METHODS OF DIAGNOSIS OF PROSTATE CANCER, COMPOSITIONS AND METHODS OF SCREENING FOR MODULATORS OF PROSTATE CANCER

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CROSS-REFERENCES TO RELATED APPLICATIONS

This application claims priority from the following applications: USSN 09/687,576 filed October 13, 2000, USSN 60/276,791 filed March 16, 2001; USSN 60/288,589, filed May 4, 2001; USSN 09/733,742, filed December 8, 2000; USSN 09/733,288, filed December 8, 2000; USSN 09/847,046, filed April 30, 2001; USSN 60/276,888, filed March 16, 2001; USSN 60/286,214, filed April 24, 2001; USSN 60/281,922, filed April 6, 2001; USSN 60/263,957, filed January 24, 2001, which are incorporated herein by reference in their entirety.

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FIELD OF THE INVENTION

The invention relates to the identification of nucleic acid and protein expression profiles and nucleic acids, products, and antibodies thereto that are involved in prostate cancer; and to the use of such expression profiles and compositions in the diagnosis, prognosis and therapy of prostate cancer. The invention further relates to methods for identifying and using agents and/or targets that inhibit prostate cancer.

BACKGROUND OF THE INVENTION

Prostate cancer is the most commonly diagnosed internal malignancy and second most common cause of cancer death in men in the U.S., resulting in approximately 40,000 deaths each year (Landis et al., *CA Cancer J. Clin.* 48:6-29 (1998); Greenlee et al., *CA Cancer J. Clin.* 50(1):7-13 (2000)), and incidence of prostate cancer has been increasing rapidly over the past 20 years in many parts of the world (Nakata et al., *Int. J. Urol.* 7(7):254-257 (2000); Majeed et al., *BJU Int.* 85(9):1058-1062 (2000)). It develops as the

result of a pathologic transformation of normal prostate cells. In tumorigenesis, the cancer cell undergoes initiation, proliferation and loss of contact inhibition, culminating in invasion of surrounding tissue and, ultimately, metastasis.

Deaths from prostate cancer are a result of metastasis of a prostate tumor. Therefore, early detection of the development of prostate cancer is critical in reducing mortality from this disease. Measuring levels of prostate-specific antigen (PSA) has become a very common method for early detection and screening, and may have contributed to the slight decrease in the mortality rate from prostate cancer in recent years (Nowroozi et al., Cancer Control 5(6):522-531 (1998)). However, many cases are not diagnosed until the disease has progressed to an advanced stage.

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Treatments such as surgery (prostatectomy), radiation therapy, and cryotherapy are potentially curative when the cancer remains localized to the prostate. Therefore, early detection of prostate cancer is important for a positive prognosis for treatment. Systemic treatment for metastatic prostate cancer is limited to hormone therapy and chemotherapy. Chemical or surgical castration has been the primary treatment for symptomatic metastatic prostate cancer for over 50 years. This testicular androgen deprivation therapy usually results in stabilization or regression of the disease (in 80% of patients), but progression of metastatic prostate cancer eventually develops (Panvichian et al., Cancer Control 3(6):493-500 (1996)). Metastatic disease is currently considered incurable, and the primary goals of treatment are to prolong survival and improve quality of life (Rago, Cancer Control 5(6):513-521 (1998)).

Thus, methods that can be used for diagnosis and prognosis of prostate cancer and effective treatment of prostate cancer, and including particularly metastatic prostate cancer, would be desirable. Accordingly, provided herein are methods that can be used in diagnosis and prognosis of prostate cancer. Further provided are methods that can be used to screen candidate bioactive agents for the ability to modulate, e.g., treat, prostate cancer. Additionally, provided herein are molecular targets and compositions for therapeutic intervention in prostate cancer and other cancers.

SUMMARY OF THE INVENTION

The present invention therefore provides nucleotide sequences of genes that are up- and down-regulated in prostate cancer cells. Such genes are useful for diagnostic purposes, and also as targets for screening for therapeutic compounds that modulate prostate cancer, such as hormones or antibodies. Other aspects of the invention will become apparent to the skilled artisan by the following description of the invention.

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In one aspect, the present invention provides a method of detecting a prostate cancer-associated transcript in a cell from a patient, the method comprising contacting a biological sample from the patient with a polynucleotide that selectively hybridizes to a sequence at least 80% identical to a sequence as shown in Tables 1-16.

In one embodiment, the present invention provides a method of determining the level of a prostate cancer associated transcript in a cell from a patient.

In one embodiment, the present invention provides a method of detecting a prostate cancer-associated transcript in a cell from a patient, the method comprising contacting a biological sample from the patient with a polynucleotide that selectively hybridizes to a sequence at least 80% identical to a sequence as shown in Tables 1-16.

In one embodiment, the polynucleotide selectively hybridizes to a sequence at least 95% identical to a sequence as shown in Tables 1-16. In another embodiment, the polynucleotide comprises a sequence as shown in Tables 1-16.

In one embodiment, the biological sample is a tissue sample. In another embodiment, the biological sample comprises isolated nucleic acids, e.g., mRNA.

In one embodiment, the polynucleotide is labeled, e.g., with a fluorescent label.

In one embodiment, the polynucleotide is immobilized on a solid surface.

In one embodiment, the patient is undergoing a therapeutic regimen to treat prostate cancer. In another embodiment, the patient is suspected of having metastatic prostate cancer.

In one embodiment, the patient is a human.

In one embodiment, the patient is suspected of having a taxol-resistant cancer.

In one embodiment, the prostate cancer associated transcript is mRNA.

In one embodiment, the method further comprises the step of amplifying nucleic acids before the step of contacting the biological sample with the polynucleotide.

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In another aspect, the present invention provides a method of monitoring the efficacy of a therapeutic treatment of prostate cancer, the method comprising the steps of: (i) providing a biological sample from a patient undergoing the therapeutic treatment; and (ii) determining the level of a prostate cancer-associated transcript in the biological sample by contacting the biological sample with a polynucleotide that selectively hybridizes to a sequence at least 80% identical to a sequence as shown in Tables 1-16, thereby monitoring the efficacy of the therapy. In a further embodiment, the patient has metastatic prostate cancer. In a further embodiment, the patient has a drug resistant (e.g., taxol resistant) form of prostate cancer.

In one embodiment, the method further comprises the step of: (iii) comparing the level of the prostate cancer-associated transcript to a level of the prostate cancer-associated transcript in a biological sample from the patient prior to, or earlier in, the therapeutic treatment.

Additionally, provided herein is a method of evaluating the effect of a candidate prostate cancer drug comprising administering the drug to a patient and removing a cell sample from the patient. The expression profile of the cell is then determined. This method may further comprise comparing the expression profile to an expression profile of a healthy individual. In a preferred embodiment, said expression profile includes a gene of Tables 1-16.

In one aspect, the present invention provides an isolated nucleic acid molecule consisting of a polynucleotide sequence as shown in Tables 1-16.

In one embodiment, an expression vector or cell comprises the isolated nucleic acid.

In one aspect, the present invention provides an isolated polypeptide which is encoded by a nucleic acid molecule having polynucleotide sequence as shown in Tables 1-16.

In another aspect, the present invention provides an antibody that specifically binds to an isolated polypeptide which is encoded by a nucleic acid molecule having polynucleotide sequence as shown in Tables 1-16.

In one embodiment, the antibody is conjugated to an effector component, e.g., a fluorescent label, a radioisotope or a cytotoxic chemical.

In one embodiment, the antibody is an antibody fragment. In another embodiment, the antibody is humanized.

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In one aspect, the present invention provides a method of detecting a prostate cancer cell in a biological sample from a patient, the method comprising contacting the biological sample with an antibody as described herein.

In another aspect, the present invention provides a method of detecting antibodies specific to prostate cancer in a patient, the method comprising contacting a biological sample from the patient with a polypeptide encoded by a nucleic acid comprising a sequence from Tables 1-16.

In another aspect, the present invention provides a method for identifying a compound that modulates a prostate cancer-associated polypeptide, the method comprising the steps of: (i) contacting the compound with a prostate cancer-associated polypeptide, the polypeptide encoded by a polynucleotide that selectively hybridizes to a sequence at least 80% identical to a sequence as shown in Tables 1-16; and (ii) determining the functional effect of the compound upon the polypeptide.

In one embodiment, the functional effect is a physical effect, an enzymatic effect, or a chemical effect.

In one embodiment, the polypeptide is expressed in a eukaryotic host cell or cell membrane. In another embodiment, the polypeptide is recombinant.

In one embodiment, the functional effect is determined by measuring ligand binding to the polypeptide.

In another aspect, the present invention provides a method of inhibiting proliferation of a prostate cancer-associated cell to treat prostate cancer in a patient, the method comprising the step of administering to the subject a therapeutically effective amount of a compound identified as described herein.

In one embodiment, the compound is an antibody.

In another aspect, the present invention provides a drug screening assay comprising the steps of: (i) administering a test compound to a mammal having prostate cancer or to a cell sample isolated therefrom; (ii) comparing the level of gene expression of a

polynucleotide that selectively hybridizes to a sequence at least 80% identical to a sequence as shown in Tables 1-16 in a treated cell or mammal with the level of gene expression of the polynucleotide in a control cell sample or mammal, wherein a test compound that modulates the level of expression of the polynucleotide is a candidate for the treatment of prostate cancer.

In one embodiment, the control is a mammal with prostate cancer or a cell sample therefrom that has not been treated with the test compound. In another embodiment, the control is a normal cell or mammal.

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In one embodiment, the test compound is administered in varying amounts or concentrations. In another embodiment, the test compound is administered for varying time periods. In another embodiment, the comparison can occur after addition or removal of the drug candidate.

In one embodiment, the levels of a plurality of polynucleotides that selectively hybridize to a sequence at least 80% identical to a sequence as shown in Tables 1-16 are individually compared to their respective levels in a control cell sample or mammal. In a preferred embodiment the plurality of polynucleotides is from three to ten.

In another aspect, the present invention provides a method for treating a mammal having prostate cancer comprising administering a compound identified by the assay described herein.

In another aspect, the present invention provides a pharmaceutical composition for treating a mammal having prostate cancer, the composition comprising a compound identified by the assay described herein and a physiologically acceptable excipient.

In one aspect, the present invention provides a method of screening drug candidates by providing a cell expressing a gene that is up- and down-regulated as in a prostate cancer. In one embodiment, a gene is selected from Tables 1-16. The method further includes adding a drug candidate to the cell and determining the effect of the drug candidate on the expression of the expression profile gene.

In one embodiment, the method of screening drug candidates includes comparing the level of expression in the absence of the drug candidate to the level of expression in the presence of the drug candidate, wherein the concentration of the drug

candidate can vary when present, and wherein the comparison can occur after addition or removal of the drug candidate. In a preferred embodiment, the cell expresses at least two expression profile genes. The profile genes may show an increase or decrease.

Also provided is a method of evaluating the effect of a candidate prostate cancer drug comprising administering the drug to a transgenic animal expressing or over-expressing the prostate cancer modulatory protein, or an animal lacking the prostate cancer modulatory protein, for example as a result of a gene knockout.

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Moreover, provided herein is a biochip comprising one or more nucleic acid segments of Tables 1-16, wherein the biochip comprises fewer than 1000 nucleic acid probes. Preferably, at least two nucleic acid segments are included. More preferably, at least three nucleic acid segments are included.

Furthermore, a method of diagnosing a disorder associated with prostate cancer is provided. The method comprises determining the expression of a gene of Tables 1-16, in a first tissue type of a first individual, and comparing the distribution to the expression of the gene from a second normal tissue type from the first individual or a second unaffected individual. A difference in the expression indicates that the first individual has a disorder associated with prostate cancer.

In a further embodiment, the biochip also includes a polynucleotide sequence of a gene that is not up- and down-regulated in prostate cancer.

In one embodiment a method for screening for a bioactive agent capable of interfering with the binding of a prostate cancer modulating protein (prostate cancer modulatory protein) or a fragment thereof and an antibody which binds to said prostate cancer modulatory protein or fragment thereof. In a preferred embodiment, the method comprises combining a prostate cancer modulatory protein or fragment thereof, a candidate bioactive agent and an antibody which binds to said prostate cancer modulatory protein or fragment thereof. The method further includes determining the binding of said prostate cancer modulatory protein or fragment thereof and said antibody. Wherein there is a change in binding, an agent is identified as an interfering agent. The interfering agent can be an agonist or an antagonist. Preferably, the agent inhibits prostate cancer.

Also provided herein are methods of eliciting an immune response in an individual. In one embodiment a method provided herein comprises administering to an

individual a composition comprising a prostate cancer modulating protein, or a fragment thereof. In another embodiment, the protein is encoded by a nucleic acid selected from those of Tables 1-16.

Further provided herein are compositions capable of eliciting an immune response in an individual. In one embodiment, a composition provided herein comprises a prostate cancer modulating protein, preferably encoded by a nucleic acid of Tables 1-16, or a fragment thereof, and a pharmaceutically acceptable carrier. In another embodiment, said composition comprises a nucleic acid comprising a sequence encoding a prostate cancer modulating protein, preferably selected from the nucleic acids of Tables 1-16, and a pharmaceutically acceptable carrier.

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Also provided are methods of neutralizing the effect of a prostate cancer protein, or a fragment thereof, comprising contacting an agent specific for said protein with said protein in an amount sufficient to effect neutralization. In another embodiment, the protein is encoded by a nucleic acid selected from those of Tables 1-16.

In another aspect of the invention, a method of treating an individual for prostate cancer is provided. In one embodiment, the method comprises administering to said individual an inhibitor of a prostate cancer modulating protein. In another embodiment, the method comprises administering to a patient having prostate cancer an antibody to a prostate cancer modulating protein conjugated to a therapeutic moiety. Such a therapeutic moiety can be a cytotoxic agent or a radioisotope.

DETAILED DESCRIPTION OF THE INVENTION

In accordance with the objects outlined above, the present invention provides novel methods for diagnosis and prognosis evaluation for prostate cancer (PC), including metastatic prostate cancer, as well as methods for screening for compositions which modulate prostate cancer. Also provided are methods for treating prostate cancer.

In addition to the other nucleic acid and peptide sequences, the present invention also relates to the identification of PAA2 as a gene that is highly over expressed in prostate cancer patient tissues. PAA2 sequence is identical to the zinc transporter ZNT4. Results presented herein demonstrate that PAA2/ZNT4 is highly expressed in prostate cancer

to accumulate zinc. Zinc uptake is regulated by prolactin and testosterone, which induce the expression of a member of the ZIP family of zinc transporters (Costello et al., 1999, J. Biol. Chem. 274:17499-17504). Zinc accumulation in the prostate functions to inhibit citrate oxidation, which results in a decrease in cellular ATP production (Costello and Franklin, 1998, Prostate 35:285-296). Cancer cells are more sensitive to decreased ATP production and have evolved to prevent zinc accumulation. Without wishing to be bound by theory, the up-regulation of ZNT4 in prostate cancer cells may result in protection of the cells from high zinc levels by its ability to pump accumulated zinc out of the cells.

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The present invention also relates to nucleic acid sequencess encoding PBH1.

PBH1 is related to human TRPC7 (transient receptor potential-related channels, NP_003298), a putative calcium channel highly expressed in brain (Nagamine et al., Genomics 54:124-131 (1998)). Trp is related to melastatin, a gene down-regulated in metastatic melanomas (Duncan et al., Cancer Res. 58:1515-1520 (1998)), and MTR1, a gene locallized to within the Beckwith-Wiedemann syndrome/Wilm's tumor susceptability region (Prawitt et al., Hum.

Mol. Genet. 9:203-216 (2000)). Without wishing to be bound by theory, it is believed that PBH1 functions as a calcium channel.

As a calcium channel, PBH1 is an ideal target for a small molecule therapeutic, or a therapeutic antibody that disrupts channel function. CD20, the target of Rituximab in non-Hodgekin's lymphoma (Maloney et al., Blood 90:2188-2195 (1997); Leget and Czuczman, Curr. Opin. Oncol. 10:548-551 (1998)), is a plasma membrane calcium channel expressed in B cells (Tedder and Engel, Immunol. Today 15:450-454 (1994)). Similarly, a small molecule, or antibody that inhibits or alters a calcium signal mediated by PBH1, will result in the death of prostate cancer cells.

PBH1, and other genes of the invention, are also be useful as targets for cytotoxic T-lymphocytes. Genes that are tumor specific, or that are expressed in immune-privileged organs, are currently being used as potential vaccine targets (Van den Eynde and Boon, Int. J. Clin. Lab. Res. 27:81-86 (1997)). The expression pattern of PBH1 indicates that it is an ideal target for cytotoxic T-lymphocytes. Thus, therapies that utilize PBH1-specific cytotoxic T-lymphocytes to induce prostate cancer cell death are also provided by this invention. See, e.g., U.S. Patent No. 6,051,227 and WO 00/32231, the disclosures of which are herein incorporated by reference.

The present invention is also related to the identification of PAA3 as a gene that is important in the modulation of prostate cancer and or breast cancer.

Tables 1-16 provide unigene cluster identification numbers, exemplar accession numbers, or genomic nucleotide position numbers for the nucleotide sequence of genes that exhibit increased or decreased expression in prostate cancer samples.

Definitions

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The term "prostate cancer protein" or "prostate cancer polynucleotide" or "prostate cancer-associated transcript" refers to nucleic acid and polypeptide polymorphic variants, alleles, mutants, and interspecies homologues that: (1) have a nucleotide sequence that has greater than about 60% nucleotide sequence identity, 65%, 70%, 75%, 80%, 85%, 90%, preferably 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98% or 99% or greater nucleotide sequence identity, preferably over a region of over a region of at least about 25, 50, 100, 200, 500, 1000, or more nucleotides, to a nucleotide sequence of or associated with a unigene cluster of Tables 1-16; (2) bind to antibodies, e.g., polyclonal antibodies, raised against an immunogen comprising an amino acid sequence encoded by a nucleotide sequence of or associated with a unigene cluster of Tables 1-16, and conservatively modified variants thereof; (3) specifically hybridize under stringent hybridization conditions to a nucleic acid sequence, or the complement thereof of Tables 1-16 and conservatively modified variants thereof or (4) have an amino acid sequence that has greater than about 60% amino acid sequence identity, 65%, 70%, 75%, 80%, 85%, 90%, preferably 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98% or 99% or greater amino sequence identity, preferably over a region of over a region of at least about 25, 50, 100, 200, 500, 1000, or more amino acid, to an amino acid sequence encoded by a nucleotide sequence of or associated with a unigene cluster of Tables 1-16. A polynucleotide or polypeptide sequence is typically from a mammal including, but not limited to, primate, e.g., human; rodent, e.g., rat, mouse, hamster; cow, pig, horse, sheep, or other mammal. A "prostate cancer polypeptide" and a "prostate cancer polynucleotide," include both naturally occurring or recombinant forms.

A "full length" prostate cancer protein or nucleic acid refers to a prostate cancer polypeptide or polynucleotide sequence, or a variant thereof, that contains all of the elements normally contained in one or more naturally occurring, wild type prostate cancer

polynucleotide or polypeptide sequences. For example, a full length prostate cancer nucleic acid will typically comprise all of the exons that encode for the full length, naturally ocurring protein. The "full length" may be prior to, or after, various stages of post-translation processing or splicing, including alternative splicing.

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"Biological sample" as used herein is a sample of biological tissue or fluid that contains nucleic acids or polypeptides, e.g., of a prostate cancer protein, polynucleotide or transcript. Such samples include, but are not limited to, tissue isolated from primates, e.g., humans, or rodents, e.g., mice, and rats. Biological samples may also include sections of tissues such as biopsy and autopsy samples, frozen sections taken for histologic purposes, blood, plasma, serum, sputum, stool, tears, mucus, hair, skin, etc. Biological samples also include explants and primary and/or transformed cell cultures derived from patient tissues. A biological sample is typically obtained from a eukaryotic organism, most preferably a mammal such as a primate e.g., chimpanzee or human; cow; dog; cat; a rodent, e.g., guinea pig, rat, mouse; rabbit; or a bird; reptile; or fish.

"Providing a biological sample" means to obtain a biological sample for use in methods described in this invention. Most often, this will be done by removing a sample of cells from an animal, but can also be accomplished by using previously isolated cells (e.g., isolated by another person, at another time, and/or for another purpose), or by performing the methods of the invention *in vivo*. Archival tissues, having treatment or outcome history, will be particularly useful.

The terms "identical" or percent "identity," in the context of two or more nucleic acids or polypeptide sequences, refer to two or more sequences or subsequences that are the same or have a specified percentage of amino acid residues or nucleotides that are the same (i.e., about 60% identity, preferably 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, or higher identity over a specified region, when compared and aligned for maximum correspondence over a comparison window or designated region) as measured using a BLAST or BLAST 2.0 sequence comparison algorithms with default parameters described below, or by manual alignment and visual inspection (see, e.g., NCBI web site http://www.ncbi.nlm.nih.gov/BLAST/ or the like). Such sequences are then said to be "substantially identical." This definition also refers to, or may be applied to, the compliment of a test sequence. The definition also includes sequences that have deletions

and/or additions, as well as those that have substitutions, as well as naturally occurring, e.g., polymorphic or allelic variants, and man-made variants. As described below, the preferred algorithms can account for gaps and the like. Preferably, identity exists over a region that is at least about 25 amino acids or nucleotides in length, or more preferably over a region that is 50-100 amino acids or nucleotides in length.

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For sequence comparison, typically one sequence acts as a reference sequence, to which test sequences are compared. When using a sequence comparison algorithm, test and reference sequences are entered into a computer, subsequence coordinates are designated, if necessary, and sequence algorithm program parameters are designated. Preferably, default program parameters can be used, or alternative parameters can be designated. The sequence comparison algorithm then calculates the percent sequence identities for the test sequences relative to the reference sequence, based on the program parameters.

A "comparison window", as used herein, includes reference to a segment of one of the number of contiguous positions selected from the group consisting typically of from 20 to 600, usually about 50 to about 200, more usually about 100 to about 150 in which a sequence may be compared to a reference sequence of the same number of contiguous positions after the two sequences are optimally aligned. Methods of alignment of sequences for comparison are well-known in the art. Optimal alignment of sequences for comparison can be conducted, e.g., by the local homology algorithm of Smith & Waterman, Adv. Appl. Math. 2:482 (1981), by the homology alignment algorithm of Needleman & Wunsch, J. Mol. Biol. 48:443 (1970), by the search for similarity method of Pearson & Lipman, Proc. Nat'l. Acad. Sci. USA 85:2444 (1988), by computerized implementations of these algorithms (GAP, BESTFIT, FASTA, and TFASTA in the Wisconsin Genetics Software Package, Genetics Computer Group, 575 Science Dr., Madison, WI), or by manual alignment and visual inspection (see, e.g., Current Protocols in Molecular Biology (Ausubel et al., eds. 1995 supplement)).

Preferred examples of algorithms that are suitable for determining percent sequence identity and sequence similarity include the BLAST and BLAST 2.0 algorithms, which are described in Altschul et al., Nuc. Acids Res. 25:3389-3402 (1977) and Altschul et al., J. Mol. Biol. 215:403-410 (1990). BLAST and BLAST 2.0 are used, with the parameters described herein, to determine percent sequence identity for the nucleic acids and proteins of

the invention. Software for performing BLAST analyses is publicly available through the National Center for Biotechnology Information (http://www.ncbi.nlm.nih.gov/). This algorithm involves first identifying high scoring sequence pairs (HSPs) by identifying short words of length W in the query sequence, which either match or satisfy some positive-valued threshold score T when aligned with a word of the same length in a database sequence. T is referred to as the neighborhood word score threshold (Altschul et al., supra). These initial neighborhood word hits act as seeds for initiating searches to find longer HSPs containing them. The word hits are extended in both directions along each sequence for as far as the cumulative alignment score can be increased. Cumulative scores are calculated using, e.g., for nucleotide sequences, the parameters M (reward score for a pair of matching residues; always > 0) and N (penalty score for mismatching residues; always < 0). For amino acid sequences, a scoring matrix is used to calculate the cumulative score. Extension of the word hits in each direction are halted when: the cumulative alignment score falls off by the quantity X from its maximum achieved value; the cumulative score goes to zero or below, due to the accumulation of one or more negative-scoring residue alignments; or the end of either sequence is reached. The BLAST algorithm parameters W, T, and X determine the sensitivity and speed of the alignment. The BLASTN program (for nucleotide sequences) uses as defaults a wordlength (W) of 11, an expectation (E) of 10, M=5, N=4 and a comparison of both strands. For amino acid sequences, the BLASTP program uses as defaults a wordlength of 3, and expectation (E) of 10, and the BLOSUM62 scoring matrix (see Henikoff & Henikoff, Proc. Natl. Acad. Sci. USA 89:10915 (1989)) alignments (B) of 50, expectation (E) of 10, M=5, N=-4, and a comparison of both strands.

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The BLAST algorithm also performs a statistical analysis of the similarity between two sequences (see, e.g., Karlin & Altschul, Proc. Nat'l. Acad. Sci. USA 90:5873-5787 (1993)). One measure of similarity provided by the BLAST algorithm is the smallest sum probability (P(N)), which provides an indication of the probability by which a match between two nucleotide or amino acid sequences would occur by chance. For example, a nucleic acid is considered similar to a reference sequence if the smallest sum probability in a comparison of the test nucleic acid to the reference nucleic acid is less than about 0.2, more preferably less than about 0.01, and most preferably less than about 0.001. Log values may be large negative numbers, e.g., 5, 10, 20, 30, 40, 40, 70, 90, 110, 150, 170, etc.

An indication that two nucleic acid sequences or polypeptides are substantially identical is that the polypeptide encoded by the first nucleic acid is immunologically cross reactive with the antibodies raised against the polypeptide encoded by the second nucleic acid, as described below. Thus, a polypeptide is typically substantially identical to a second polypeptide, e.g., where the two peptides differ only by conservative substitutions. Another indication that two nucleic acid sequences are substantially identical is that the two molecules or their complements hybridize to each other under stringent conditions, as described below. Yet another indication that two nucleic acid sequences are substantially identical is that the same primers can be used to amplify the sequences.

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A "host cell" is a naturally occurring cell or a transformed cell that contains an expression vector and supports the replication or expression of the expression vector. Host cells may be cultured cells, explants, cells *in vivo*, and the like. Host cells may be prokaryotic cells such as *E. coli*, or eukaryotic cells such as yeast, insect, amphibian, or mammalian cells such as CHO, HeLa, and the like (*see, e.g.*, the American Type Culture Collection catalog or web site, www.atcc.org).

The terms "isolated," "purified," or "biologically pure" refer to material that is substantially or essentially free from components that normally accompany it as found in its native state. Purity and homogeneity are typically determined using analytical chemistry techniques such as polyacrylamide gel electrophoresis or high performance liquid chromatography. A protein or nucleic acid that is the predominant species present in a preparation is substantially purified. In particular, an isolated nucleic acid is separated from some open reading frames that naturally flank the gene and encode proteins other than protein encoded by the gene. The term "purified" in some embodiments denotes that a nucleic acid or protein gives rise to essentially one band in an electrophoretic gel. Preferably, it means that the nucleic acid or protein is at least 85% pure, more preferably at least 95% pure, and most preferably at least 99% pure. "Purify" or "purification" in other embodiments means removing at least one contaminant from the composition to be purified. In this sense, purification does not require that the purified compound be homogenous, e.g., 100% pure.

The terms "polypeptide," "peptide" and "protein" are used interchangeably herein to refer to a polymer of amino acid residues. The terms apply to amino acid polymers in which one or more amino acid residue is an artificial chemical mimetic of a corresponding

naturally occurring amino acid, as well as to naturally occurring amino acid polymers, those containing modified residues, and non-naturally occurring amino acid polymer.

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The term "amino acid" refers to naturally occurring and synthetic amino acids, as well as amino acid analogs and amino acid mimetics that function similarly to the naturally occurring amino acids. Naturally occurring amino acids are those encoded by the genetic code, as well as those amino acids that are later modified, e.g., hydroxyproline, γ-carboxyglutamate, and O-phosphoserine. Amino acid analogs refers to compounds that have the same basic chemical structure as a naturally occurring amino acid, e.g., an α carbon that is bound to a hydrogen, a carboxyl group, an amino group, and an R group, e.g., homoserine, norleucine, methionine sulfoxide, methionine methyl sulfonium. Such analogs may have modified R groups (e.g., norleucine) or modified peptide backbones, but retain the same basic chemical structure as a naturally occurring amino acid. Amino acid mimetics refers to chemical compounds that have a structure that is different from the general chemical structure of an amino acid, but that functions similarly to a naturally occurring amino acid.

Amino acids may be referred to herein by either their commonly known three letter symbols or by the one-letter symbols recommended by the IUPAC-IUB Biochemical Nomenclature Commission. Nucleotides, likewise, may be referred to by their commonly accepted single-letter codes.

"Conservatively modified variants" applies to both amino acid and nucleic acid sequences. With respect to particular nucleic acid sequences, conservatively modified variants refers to those nucleic acids which encode identical or essentially identical amino acid sequences, or where the nucleic acid does not encode an amino acid sequence, to essentially identical or associated, e.g., naturally contiguous, sequences. Because of the degeneracy of the genetic code, a large number of functionally identical nucleic acids encode most proteins. For instance, the codons GCA, GCC, GCG and GCU all encode the amino acid alanine. Thus, at every position where an alanine is specified by a codon, the codon can be altered to another of the corresponding codons described without altering the encoded polypeptide. Such nucleic acid variations are "silent variations," which are one species of conservatively modified variations. Every nucleic acid sequence herein which encodes a polypeptide also describes silent variations of the nucleic acid. One of skill will recognize that in certain contexts each codon in a nucleic acid (except AUG, which is ordinarily the

only codon for methionine, and TGG, which is ordinarily the only codon for tryptophan) can be modified to yield a functionally identical molecule. Accordingly, often silent variations of a nucleic acid which encodes a polypeptide is implicit in a described sequence with respect to the expression product, but not with respect to actual probe sequences.

As to amino acid sequences, one of skill will recognize that individual substitutions, deletions or additions to a nucleic acid, peptide, polypeptide, or protein sequence which alters, adds or deletes a single amino acid or a small percentage of amino acids in the encoded sequence is a "conservatively modified variant" where the alteration results in the substitution of an amino acid with a chemically similar amino acid.

Conservative substitution tables providing functionally similar amino acids are well known in the art. Such conservatively modified variants are in addition to and do not exclude polymorphic variants, interspecies homologs, and alleles of the invention typically conservative substitutions for one another: 1) Alanine (A), Glycine (G); 2) Aspartic acid (D), Glutamic acid (E); 3) Asparagine (N), Glutamine (Q); 4) Arginine (R), Lysine (K); 5)

Isoleucine (I), Leucine (L), Methionine (M), Valine (V); 6) Phenylalanine (F), Tyrosine (Y), Tryptophan (W); 7) Serine (S), Threonine (T); and 8) Cysteine (C), Methionine (M) (see, e.g., Creighton, Proteins (1984)).

Macromolecular structures such as polypeptide structures can be described in terms of various levels of organization. For a general discussion of this organization, see, e.g., Alberts et al., Molecular Biology of the Cell (3rd ed., 1994) and Cantor & Schimmel, Biophysical Chemistry Part I: The Conformation of Biological Macromolecules (1980). "Primary structure" refers to the amino acid sequence of a particular peptide. "Secondary structure" refers to locally ordered, three dimensional structures within a polypeptide. These structures are commonly known as domains. Domains are portions of a polypeptide that often form a compact unit of the polypeptide and are typically 25 to approximately 500 amino acids long. Typical domains are made up of sections of lesser organization such as stretches of β -sheet and α -helices. "Tertiary structure" refers to the complete three dimensional structure of a polypeptide monomer. "Quaternary structure" refers to the three dimensional structure formed, usually by the noncovalent association of independent tertiary units. Anisotropic terms are also known as energy terms.

"Nucleic acid" or "oligonucleotide" or "polynucleotide" or grammatical equivalents used herein means at least two nucleotides covalently linked together. Oligonucleotides are typically from about 5, 6, 7, 8, 9, 10, 12, 15, 25, 30, 40, 50 or more nucleotides in length, up to about 100 nucleotides in length. Nucleic acids and polynucleotides are a polymers of any length, including longer lengths, e.g., 200, 300, 500, 1000, 2000, 3000, 5000, 7000, 10,000, etc. A nucleic acid of the present invention will generally contain phosphodiester bonds, although in some cases, nucleic acid analogs are included that may have alternate backbones, comprising, e.g., phosphoramidate, phosphorothioate, phosphorodithioate, or O-methylphophoroamidite linkages (see Eckstein, Oligonucleotides and Analogues: A Practical Approach, Oxford University Press); and peptide nucleic acid backbones and linkages. Other analog nucleic acids include those with positive backbones; non-ionic backbones, and non-ribose backbones, including those described in U.S. Patent Nos. 5,235,033 and 5,034,506, and Chapters 6 and 7, ASC Symposium Series 580, Carbohydrate Modifications in Antisense Research, Sanghui & Cook, eds.. Nucleic acids containing one or more carbocyclic sugars are also included within one definition of nucleic acids. Modifications of the ribose-phosphate backbone may be done for a variety of reasons, e.g. to increase the stability and half-life of such molecules in physiological environments or as probes on a biochip. Mixtures of naturally occurring nucleic acids and analogs can be made; alternatively, mixtures of different nucleic acid analogs, and mixtures of naturally occurring nucleic acids and analogs may be made.

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A variety of references disclose such nucleic acid analogs, including, for example, phosphoramidate (Beaucage et al., Tetrahedron 49(10):1925 (1993) and references therein; Letsinger, J. Org. Chem. 35:3800 (1970); Sprinzl et al., Eur. J. Biochem. 81:579 (1977); Letsinger et al., Nucl. Acids Res. 14:3487 (1986); Sawai et al, Chem. Lett. 805 (1984), Letsinger et al., J. Am. Chem. Soc. 110:4470 (1988); and Pauwels et al., Chemica Scripta 26:141 91986)), phosphorothioate (Mag et al., Nucleic Acids Res. 19:1437 (1991); and U.S. Patent No. 5,644,048), phosphorodithioate (Briu et al., J. Am. Chem. Soc. 111:2321 (1989), O-methylphophoroamidite linkages (see Eckstein, Oligonucleotides and Analogues: A Practical Approach, Oxford University Press), and peptide nucleic acid backbones and linkages (see Egholm, J. Am. Chem. Soc. 114:1895 (1992); Meier et al., Chem. Int. Ed. Engl. 31:1008 (1992); Nielsen, Nature, 365:566 (1993); Carlsson et al., Nature 380:207 (1996), all

of which are incorporated by reference). Other analog nucleic acids include those with positive backbones (Denpcy et al., Proc. Natl. Acad. Sci. USA 92:6097 (1995); non-ionic backbones (U.S. Patent Nos. 5,386,023, 5,637,684, 5,602,240, 5,216,141 and 4,469,863; Kiedrowshi et al., Angew. Chem. Intl. Ed. English 30:423 (1991); Letsinger et al., J. Am. 5 Chem. Soc. 110:4470 (1988); Letsinger et al., Nucleoside & Nucleotide 13:1597 (1994); Chapters 2 and 3, ASC Symposium Series 580, "Carbohydrate Modifications in Antisense Research", Ed. Y.S. Sanghui and P. Dan Cook; Mesmaeker et al., Bioorganic & Medicinal Chem. Lett. 4:395 (1994); Jeffs et al., J. Biomolecular NMR 34:17 (1994); Tetrahedron Lett. 37:743 (1996)) and non-ribose backbones, including those described in U.S. Patent Nos. 10 5,235,033 and 5,034,506, and Chapters 6 and 7, ASC Symposium Series 580, "Carbohydrate Modifications in Antisense Research", Ed. Y.S. Sanghui and P. Dan Cook. Nucleic acids containing one or more carbocyclic sugars are also included within one definition of nucleic acids (see Jenkins et al., Chem. Soc. Rev. (1995) pp 169-176). Several nucleic acid analogs are described in Rawls, C & E News June 2, 1997 page 35. All of these references are hereby 15 expressly incorporated by reference.

Particularly preferred are peptide nucleic acids (PNA) which includes peptide nucleic acid analogs. These backbones are substantially non-ionic under neutral conditions, in contrast to the highly charged phosphodiester backbone of naturally occurring nucleic acids. This results in two advantages. First, the PNA backbone exhibits improved hybridization kinetics. PNAs have larger changes in the melting temperature (T_m) for mismatched versus perfectly matched basepairs. DNA and RNA typically exhibit a 2-4°C drop in T_m for an internal mismatch. With the non-ionic PNA backbone, the drop is closer to 7-9°C. Similarly, due to their non-ionic nature, hybridization of the bases attached to these backbones is relatively insensitive to salt concentration. In addition, PNAs are not degraded by cellular enzymes, and thus can be more stable.

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The nucleic acids may be single stranded or double stranded, as specified, or contain portions of both double stranded or single stranded sequence. As will be appreciated by those in the art, the depiction of a single strand also defines the sequence of the complementary strand; thus the sequences described herein also provide the complement of the sequence. The nucleic acid may be DNA, both genomic and cDNA, RNA or a hybrid, where the nucleic acid may contain combinations of deoxyribo- and ribo-nucleotides, and

combinations of bases, including uracil, adenine, thymine, cytosine, guanine, inosine, xanthine hypoxanthine, isocytosine, isoguanine, etc. "Transcript" typically refers to a naturally occurring RNA, e.g., a pre-mRNA, hnRNA, or mRNA. As used herein, the term "nucleoside" includes nucleotides and nucleoside and nucleotide analogs, and modified nucleosides such as amino modified nucleosides. In addition, "nucleoside" includes non-naturally occurring analog structures. Thus, e.g. the individual units of a peptide nucleic acid, each containing a base, are referred to herein as a nucleoside.

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A "label" or a "detectable moiety" is a composition detectable by spectroscopic, photochemical, biochemical, immunochemical, chemical, or other physical means. For example, useful labels include fluorescent dyes, electron-dense reagents, enzymes (e.g., as commonly used in an ELISA), biotin, digoxigenin, or haptens and proteins or other entities which can be made detectable, e.g., by incorporating a radiolabel into the peptide or used to detect antibodies specifically reactive with the peptide. The radioisotope may be, for example, 3H, 14C, 32P, 35S, or 125I. In some cases, particularly using antibodies against the proteins of the invention, the radioisotopes are used as toxic moieties, as described below. The labels may be incorporated into the prostate cancer nucleic acids, proteins and antibodies at any position. Any method known in the art for conjugating the antibody to the label may be employed, including those methods described by Hunter et al., Nature, 144:945 (1962); David et al., Biochemistry, 13:1014 (1974); Pain et al., J. Immunol. Meth., 40:219 (1981); and Nygren, J. Histochem. and Cytochem., 30:407 (1982). The lifetime of radiolabeled peptides or radiolabeled antibody compositions may extended by the addition of substances that stablize the radiolabeled peptide or antibody and protect it from degradation. Any substance or combination of substances that stablize the radiolabeled peptide or antibody may be used including those substances disclosed in US Patent No. 5,961,955.

An "effector" or "effector moiety" or "effector component" is a molecule that is bound (or linked, or conjugated), either covalently, through a linker or a chemical bond, or noncovalently, through ionic, van der Waals, electrostatic, or hydrogen bonds, to an antibody. The "effector" can be a variety of molecules including, e.g., detection moieties including radioactive compounds, fluorescent compounds, an enzyme or substrate, tags such as epitope tags, a toxin; activatable moieties, a chemotherapeutic agent; a lipase; an antibiotic; or a radioisotope emitting "hard" e.g., beta radiation.

A "labeled nucleic acid probe or oligonucleotide" is one that is bound, either covalently, through a linker or a chemical bond, or noncovalently, through ionic, van der Waals, electrostatic, or hydrogen bonds to a label such that the presence of the probe may be detected by detecting the presence of the label bound to the probe. Alternatively, method using high affinity interactions may achieve the same results where one of a pair of binding partners binds to the other, e.g., biotin, streptavidin.

As used herein a "nucleic acid probe or oligonucleotide" is defined as a nucleic acid capable of binding to a target nucleic acid of complementary sequence through one or more types of chemical bonds, usually through complementary base pairing, usually through hydrogen bond formation. As used herein, a probe may include natural (i.e., A, G, C, or T) or modified bases (7-deazaguanosine, inosine, etc.). In addition, the bases in a probe may be joined by a linkage other than a phosphodiester bond, so long as it does not functionally interfere with hybridization. Thus, e.g., probes may be peptide nucleic acids in which the constituent bases are joined by peptide bonds rather than phosphodiester linkages. It will be understood by one of skill in the art that probes may bind target sequences lacking complete complementarity with the probe sequence depending upon the stringency of the hybridization conditions. The probes are preferably directly labeled as with isotopes, chromophores, lumiphores, chromogens, or indirectly labeled such as with biotin to which a streptavidin complex may later bind. By assaying for the presence or absence of the probe, one can detect the presence or absence of the select sequence or subsequence. Diagnosis or prognosis may be based at the genomic level, or at the level of RNA or protein expression.

The term "recombinant" when used with reference, e.g., to a cell, or nucleic acid, protein, or vector, indicates that the cell, nucleic acid, protein or vector, has been modified by the introduction of a heterologous nucleic acid or protein or the alteration of a native nucleic acid or protein, or that the cell is derived from a cell so modified. Thus, e.g., recombinant cells express genes that are not found within the native (non-recombinant) form of the cell or express native genes that are otherwise abnormally expressed, under expressed or not expressed at all. By the term "recombinant nucleic acid" herein is meant nucleic acid, originally formed *in vitro*, in general, by the manipulation of nucleic acid, e.g., using polymerases and endonucleases, in a form not normally found in nature. In this manner, operably linkage of different sequences is achieved. Thus an isolated nucleic acid, in a linear

form, or an expression vector formed *in vitro* by ligating DNA molecules that are not normally joined, are both considered recombinant for the purposes of this invention. It is understood that once a recombinant nucleic acid is made and reintroduced into a host cell or organism, it will replicate non-recombinantly, i.e., using the *in vivo* cellular machinery of the host cell rather than *in vitro* manipulations; however, such nucleic acids, once produced recombinantly, although subsequently replicated non-recombinantly, are still considered recombinant for the purposes of the invention. Similarly, a "recombinant protein" is a protein made using recombinant techniques, i.e., through the expression of a recombinant nucleic acid as depicted above.

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The term "heterologous" when used with reference to portions of a nucleic acid indicates that the nucleic acid comprises two or more subsequences that are not normally found in the same relationship to each other in nature. For instance, the nucleic acid is typically recombinantly produced, having two or more sequences, e.g., from unrelated genes arranged to make a new functional nucleic acid, e.g., a promoter from one source and a coding region from another source. Similarly, a heterologous protein will often refer to two or more subsequences that are not found in the same relationship to each other in nature (e.g., a fusion protein).

A "promoter" is defined as an array of nucleic acid control sequences that direct transcription of a nucleic acid. As used herein, a promoter includes necessary nucleic acid sequences near the start site of transcription, such as, in the case of a polymerase II type promoter, a TATA element. A promoter also optionally includes distal enhancer or repressor elements, which can be located as much as several thousand base pairs from the start site of transcription. A "constitutive" promoter is a promoter that is active under most environmental and developmental conditions. An "inducible" promoter is a promoter that is active under environmental or developmental regulation. The term "operably linked" refers to a functional linkage between a nucleic acid expression control sequence (such as a promoter, or array of transcription factor binding sites) and a second nucleic acid sequence, wherein the expression control sequence directs transcription of the nucleic acid corresponding to the second sequence.

An "expression vector" is a nucleic acid construct, generated recombinantly or synthetically, with a series of specified nucleic acid elements that permit transcription of a

particular nucleic acid in a host cell. The expression vector can be part of a plasmid, virus, or nucleic acid fragment. Typically, the expression vector includes a nucleic acid to be transcribed operably linked to a promoter.

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The phrase "selectively (or specifically) hybridizes to refers to the binding, duplexing, or hybridizing of a molecule only to a particular nucleotide sequence that is determinative of the presence of the nucleotide sequence, in a heterogeneous population of nucleic acids and other biologics (e.g., total cellular or library DNA or RNA). Similarly, the phrase "specifically (or selectively) binds" to an antibody or "specifically (or selectively) immunoreactive with," when referring to a protein or peptide, refers to a binding reaction that is determinative of the presence of the protein, in a heterogeneous population of proteins and other biologics. Thus, under designated immunoassay or nucleic acid hybridization conditions, the specified antibodies or nucleic acid probes bind to a particular protein nucleotide sequences at least two times the background and more typically more than 10 to 100 times background.

Specific binding to an antibody under such conditions requires an antibody that is selected for its specificity for a particular protein. For example, polyclonal antibodies raised to a particular protein, polymorphic variants, alleles, orthologs, and conservatively modified variants, or splice variants, or portions thereof, can be selected to obtain only those polyclonal antibodies that are specifically immunoreactive with the desired prostact cancer protein and not with other proteins. This selection may be achieved by subtracting out antibodies that cross-react with other molecules. A variety of immunoassay formats may be used to select antibodies specifically immunoreactive with a particular protein. For example, solid-phase ELISA immunoassays are routinely used to select antibodies specifically immunoreactive with a protein (see, e.g., Harlow & Lane, Antibodies, A Laboratory Manual (1988) for a description of immunoassay formats and conditions that can be used to determine specific immunoreactivity).

The phrase "stringent hybridization conditions" refers to conditions under which a probe will hybridize to its target subsequence, typically in a complex mixture of nucleic acids, but to no other sequences. Stringent conditions are sequence-dependent and will be different in different circumstances. Longer sequences hybridize specifically at higher temperatures. An extensive guide to the hybridization of nucleic acids is found in

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Tijssen, Techniques in Biochemistry and Molecular Biology--Hybridization with Nucleic Probes, "Overview of principles of hybridization and the strategy of nucleic acid assays" (1993). Generally, stringent conditions are selected to be about 5-10°C lower than the thermal melting point (T_m) for the specific sequence at a defined ionic strength pH. The T_m is the temperature (under defined ionic strength, pH, and nucleic concentration) at which 50% of the probes complementary to the target hybridize to the target sequence at equilibrium (as the target sequences are present in excess, at T_m, 50% of the probes are occupied at equilibrium). Stringent conditions will be those in which the salt concentration is less than about 1.0 M sodium ion, typically about 0.01 to 1.0 M sodium ion concentration (or other salts) at pH 7.0 to 8.3 and the temperature is at least about 30°C for short probes (e.g., 10 to 50 nucleotides) and at least about 60°C for long probes (e.g., greater than 50 nucleotides). Stringent conditions may also be achieved with the addition of destabilizing agents such as formamide. For selective or specific hybridization, a positive signal is at least two times background; preferably 10 times background hybridization. Exemplary stringent hybridization conditions can be as following: 50% formamide, 5x SSC, and 1% SDS, incubating at 42°C, or, 5x SSC, 1% SDS, incubating at 65°C, with wash in 0.2x SSC, and 0.1% SDS at 65°C. For PCR, a temperature of about 36°C is typical for low stringency amplification, although annealing temperatures may vary between about 32°C and 48°C depending on primer length. For high stringency PCR amplification, a temperature of about 62°C is typical, although high stringency annealing temperatures can range from about 50°C to about 65°C, depending on the primer length and specificity. Typical cycle conditions for both high and low stringency amplifications include a denaturation phase of 90°C - 95°C for 30 sec - 2 min., an annealing phase lasting 30 sec. - 2 min., and an extension phase of about 72°C for 1 - 2 min. Protocols and guidelines for low and high stringency amplification reactions are provided, e.g., in Innis et al. (1990) PCR Protocols, A Guide to Methods and Applications, Academic Press, Inc. N.Y.).

Nucleic acids that do not hybridize to each other under stringent conditions are still substantially identical if the polypeptides which they encode are substantially identical. This occurs, e.g., when a copy of a nucleic acid is created using the maximum codon degeneracy permitted by the genetic code. In such cases, the nucleic acids typically hybridize

under moderately stringent hybridization conditions. Exemplary "moderately stringent hybridization conditions" include a hybridization in a buffer of 40% formamide, 1 M NaCl, 1% SDS at 37°C, and a wash in 1X SSC at 45°C. A positive hybridization is at least twice background. Those of ordinary skill will readily recognize that alternative hybridization and wash conditions can be utilized to provide conditions of similar stringency. Additional guidelines for determining hybridization parameters are provided in numerous reference, e.g., and Current Protocols in Molecular Biology, ed. Ausubel, et al.

The phrase "functional effects" in the context of assays for testing compounds that modulate activity of a prostate cancer protein includes the determination of a parameter that is indirectly or directly under the influence of the prostate cancer protein or nucleic acid, e.g., a functional, physical, or chemical effect, such as the ability to decrease prostate cancer. It includes ligand binding activity; cell growth on soft agar; anchorage dependence; contact inhibition and density limitation of growth; cellular proliferation; cellular transformation; growth factor or serum dependence; tumor specific marker levels; invasiveness into Matrigel; tumor growth and metastasis *in vivo*; mRNA and protein expression in cells undergoing metastasis, and other characteristics of prostate cancer cells. "Functional effects" include *in vitro*, *in vivo*, and *ex vivo* activities.

By "determining the functional effect" is meant assaying for a compound that increases or decreases a parameter that is indirectly or directly under the influence of a prostate cancer protein sequence, e.g., functional, enzymatic, physical and chemical effects. Such functional effects can be measured by any means known to those skilled in the art, e.g., changes in spectroscopic characteristics (e.g., fluorescence, absorbance, refractive index), hydrodynamic (e.g., shape), chromatographic, or solubility properties for the protein, measuring inducible markers or transcriptional activation of the prostate cancer protein; measuring binding activity or binding assays, e.g. binding to antibodies or other ligands, and measuring cellular proliferation. Determination of the functional effect of a compound on prostate cancer can also be performed using prostate cancer assays known to those of skill in the art such as an *in vitro* assays, e.g., cell growth on soft agar; anchorage dependence; contact inhibition and density limitation of growth; cellular proliferation; cellular transformation; growth factor or serum dependence; tumor specific marker levels; invasiveness into Matrigel; tumor growth and metastasis *in vivo*; mRNA and protein

expression in cells undergoing metastasis, and other characteristics of prostate cancer cells. The functional effects can be evaluated by many means known to those skilled in the art, e.g., microscopy for quantitative or qualitative measures of alterations in morphological features, measurement of changes in RNA or protein levels for prostate cancer-associated sequences, measurement of RNA stability, identification of downstream or reporter gene expression (CAT, luciferase, β -gal, GFP and the like), e.g., via chemiluminescence, fluorescence, colorimetric reactions, antibody binding, inducible markers, and ligand binding assays.

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"Inhibitors", "activators", and "modulators" of prostate cancer polynucleotide and polypeptide sequences are used to refer to activating, inhibitory, or modulating molecules or compounds identified using in vitro and in vivo assays of prostate cancer polynucleotide and polypeptide sequences. Inhibitors are compounds that, e.g., bind to, partially or totally block activity, decrease, prevent, delay activation, inactivate, desensitize, or down regulate the activity or expression of prostate cancer proteins, e.g., antagonists. Antisense nucleic acids may seem to inhibit expression and subsequent function of the protein. "Activators" are compounds that increase, open, activate, facilitate, enhance activation, sensitize, agonize, or up regulate prostate cancer protein activity. Inhibitors, activators, or modulators also include genetically modified versions of prostate cancer proteins, e.g., versions with altered activity, as well as naturally occurring and synthetic ligands, antagonists, agonists, antibodies, small chemical molecules and the like. Such assays for inhibitors and activators include, e.g., expressing the prostate cancer protein in vitro, in cells, or cell membranes, applying putative modulator compounds, and then determining the functional effects on activity, as described above. Activators and inhibitors of prostate cancer can also be identified by incubating prostate cancer cells with the test compound and determining increases or decreases in the expression of 1 or more prostate cancer proteins, e.g., 1, 2, 3, 4, 5, 10, 15, 20, 25, 30, 40, 50 or more prostate cancer proteins, such as prostate cancer proteins encoded by the sequences set out in Tables 1-16.

Samples or assays comprising prostate cancer proteins that are treated with a potential activator, inhibitor, or modulator are compared to control samples without the inhibitor, activator, or modulator to examine the extent of inhibition. Control samples (untreated with inhibitors) are assigned a relative protein activity value of 100%. Inhibition of a polypeptide is achieved when the activity value relative to the control is about 80%,

preferably 50%, more preferably 25-0%. Activation of a prostate cancer polypeptide is achieved when the activity value relative to the control (untreated with activators) is 110%, more preferably 150%, more preferably 200-500% (i.e., two to five fold higher relative to the control), more preferably 1000-3000% higher.

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The phrase "changes in cell growth" refers to any change in cell growth and proliferation characteristics *in vitro* or *in vivo*, such as formation of foci, anchorage independence, semi-solid or soft agar growth, changes in contact inhibition and density limitation of growth, loss of growth factor or serum requirements, changes in cell morphology, gaining or losing immortalization, gaining or losing tumor specific markers, ability to form or suppress tumors when injected into suitable animal hosts, and/or immortalization of the cell. *See, e.g.*, Freshney, *Culture of Animal Cells a Manual of Basic Technique* pp. 231-241 (3rd ed. 1994).

"Tumor cell" refers to precancerous, cancerous, and normal cells in a tumor.

"Cancer cells," "transformed" cells or "transformation" in tissue culture, refers to spontaneous or induced phenotypic changes that do not necessarily involve the uptake of new genetic material. Although transformation can arise from infection with a transforming virus and incorporation of new genomic DNA, or uptake of exogenous DNA, it can also arise spontaneously or following exposure to a carcinogen, thereby mutating an endogenous gene. Transformation is associated with phenotypic changes, such as immortalization of cells, aberrant growth control, nonmorphological changes, and/or malignancy (see, Freshney, Culture of Animal Cells a Manual of Basic Technique (3rd ed. 1994)).

"Antibody" refers to a polypeptide comprising a framework region from an immunoglobulin gene or fragments thereof that specifically binds and recognizes an antigen. The recognized immunoglobulin genes include the kappa, lambda, alpha, gamma, delta, epsilon, and mu constant region genes, as well as the myriad immunoglobulin variable region genes. Light chains are classified as either kappa or lambda. Heavy chains are classified as gamma, mu, alpha, delta, or epsilon, which in turn define the immunoglobulin classes, IgG, IgM, IgA, IgD and IgE, respectively. Typically, the antigen-binding region of an antibody or its functional equivalent will be most critical in specificity and affinity of binding. See Paul, Fundamental Immunology.

An exemplary immunoglobulin (antibody) structural unit comprises a tetramer. Each tetramer is composed of two identical pairs of polypeptide chains, each pair having one "light" (about 25 kD) and one "heavy" chain (about 50-70 kD). The N-terminus of each chain defines a variable region of about 100 to 110 or more amino acids primarily responsible for antigen recognition. The terms variable light chain (V_L) and variable heavy chain (V_H) refer to these light and heavy chains respectively.

Antibodies exist, e.g., as intact immunoglobulins or as a number of well-characterized fragments produced by digestion with various peptidases. Thus, e.g., pepsin digests an antibody below the disulfide linkages in the hinge region to produce F(ab)'₂, a dimer of Fab which itself is a light chain joined to V_H-C_H1 by a disulfide bond. The F(ab)'₂ may be reduced under mild conditions to break the disulfide linkage in the hinge region, thereby converting the F(ab)'₂ dimer into an Fab' monomer. The Fab' monomer is essentially Fab with part of the hinge region (see Fundamental Immunology (Paul ed., 3d ed. 1993). While various antibody fragments are defined in terms of the digestion of an intact antibody, one of skill will appreciate that such fragments may be synthesized de novo either chemically or by using recombinant DNA methodology. Thus, the term antibody, as used herein, also includes antibody fragments either produced by the modification of whole antibodies, or those synthesized de novo using recombinant DNA methodologies (e.g., single chain Fv) or those identified using phage display libraries (see, e.g., McCafferty et al., Nature 348:552-554 (1990))

For preparation of antibodies, e.g., recombinant, monoclonal, or polyclonal antibodies, many technique known in the art can be used (see, e.g., Kohler & Milstein, Nature 256:495-497 (1975); Kozbor et al., Immunology Today 4:72 (1983); Cole et al., pp. 77-96 in Monoclonal Antibodies and Cancer Therapy (1985); Coligan, Current Protocols in Immunology (1991); Harlow & Lane, Antibodies, A Laboratory Manual (1988); and Goding, Monoclonal Antibodies: Principles and Practice (2d ed. 1986)). Techniques for the production of single chain antibodies (U.S. Patent 4,946,778) can be adapted to produce antibodies to polypeptides of this invention. Also, transgenic mice, or other organisms such as other mammals, may be used to express humanized antibodies. Alternatively, phage display technology can be used to identify antibodies and heteromeric Fab fragments that

specifically bind to selected antigens (see, e.g., McCafferty et al., Nature 348:552-554 (1990); Marks et al., Biotechnology 10:779-783 (1992)).

A "chimeric antibody" is an antibody molecule in which (a) the constant region, or a portion thereof, is altered, replaced or exchanged so that the antigen binding site (variable region) is linked to a constant region of a different or altered class, effector function and/or species, or an entirely different molecule which confers new properties to the chimeric antibody, *e.g.*, an enzyme, toxin, hormone, growth factor, drug, etc.; or (b) the variable region, or a portion thereof, is altered, replaced or exchanged with a variable region having a different or altered antigen specificity.

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Identification of prostate cancer-associated sequences

In one aspect, the expression levels of genes are determined in different patient samples for which diagnosis information is desired, to provide expression profiles. An expression profile of a particular sample is essentially a "fingerprint" of the state of the sample; while two states may have any particular gene similarly expressed, the evaluation of a number of genes simultaneously allows the generation of a gene expression profile that is characteristic of the state of the cell. That is, normal tissue (e.g., normal prostate or other tissue) may be distinguished from cancerous or metastatic cancerous tissue of the prostate, or prostate cancer tissue or metastatic prostate cancerous tissue can be compared with tissue samples of prostate and other tissues from surviving cancer patients. By comparing expression profiles of tissue in known different prostate cancer states, information regarding which genes are important (including both up- and down-regulation of genes) in each of these states is obtained.

The identification of sequences that are differentially expressed in prostate cancer versus non-prostate cancer tissue allows the use of this information in a number of ways. For example, a particular treatment regime may be evaluated: does a chemotherapeutic drug act to down-regulate prostate cancer, and thus tumor growth or recurrence, in a particular patient. Similarly, diagnosis and treatment outcomes may be done or confirmed by comparing patient samples with the known expression profiles. Metastatic tissue can also be analyzed to determine the stage of prostate cancer in the tissue. Furthermore, these gene expression profiles (or individual genes) allow screening of drug candidates with an eye to

mimicking or altering a particular expression profile; e.g., screening can be done for drugs that suppress the prostate cancer expression profile. This may be done by making biochips comprising sets of the important prostate cancer genes, which can then be used in these screens. These methods can also be done on the protein basis; that is, protein expression levels of the prostate cancer proteins can be evaluated for diagnostic purposes or to screen candidate agents. In addition, the prostate cancer nucleic acid sequences can be administered for gene therapy purposes, including the administration of antisense nucleic acids, or the prostate cancer proteins (including antibodies and other modulators thereof) administered as therapeutic drugs.

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Thus the present invention provides nucleic acid and protein sequences that are differentially expressed in prostate cancer, herein termed "prostate cancer sequences." As outlined below, prostate cancer sequences include those that are up-regulated (i.e., expressed at a higher level) in prostate cancer, as well as those that are down-regulated (i.e., expressed at a lower level). In a preferred embodiment, the prostate cancer sequences are from humans; however, as will be appreciated by those in the art, prostate cancer sequences from other organisms may be useful in animal models of disease and drug evaluation; thus, other prostate cancer sequences are provided, from vertebrates, including mammals, including rodents (rats, mice, hamsters, guinea pigs, etc.), primates, farm animals (including sheep, goats, pigs, cows, horses, etc.) and pets, e.g., (dogs, cats, etc.). Prostate cancer sequences from other organisms may be obtained using the techniques outlined below.

Prostate cancer sequences can include both nucleic acid and amino acid sequences. As will be appreciated by those in the art and is more fully outlined below, prostate cancer nucleic acid sequences are useful in a variety of applications, including diagnostic applications, which will detect naturally occurring nucleic acids, as well as screening applications; e.g., biochips comprising nucleic acid probes or PCR microtiter plates with selected probes to the prostate cancer sequences can be generated.

A prostate cancer sequence can be initially identified by substantial nucleic acid and/or amino acid sequence homology to the prostate cancer sequences outlined herein. Such homology can be based upon the overall nucleic acid or amino acid sequence, and is generally determined as outlined below, using either homology programs or hybridization conditions.

For identifying prostate cancer-associated sequences, the prostate cancer screen typically includes comparing genes identified in different tissues, e.g., normal and cancerous tissues, or tumor tissue samples from patients who have metastatic disease vs. non metastatic tissue. Other suitable tissue comparisons include comparing prostate cancer samples with metastatic cancer samples from other cancers, such as lung, breast, gastrointestinal cancers, ovarian, etc. Samples of different stages of prostate cancer, e.g., survivor tissue, drug resistant states, and tissue undergoing metastasis, are applied to biochips comprising nucleic acid probes. The samples are first microdissected, if applicable, and treated as is known in the art for the preparation of mRNA. Suitable biochips are commercially available, e.g. from Affymetrix. Gene expression profiles as described herein are generated and the data analyzed.

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In one embodiment, the genes showing changes in expression as between normal and disease states are compared to genes expressed in other normal tissues, preferably normal prostate, but also including, and not limited to lung, heart, brain, liver, breast, kidney, muscle, colon, small intestine, large intestine, spleen, bone and placenta. In a preferred embodiment, those genes identified during the prostate cancer screen that are expressed in any significant amount in other tissues are removed from the profile, although in some embodiments, this is not necessary. That is, when screening for drugs, it is usually preferable that the target be disease specific, to minimize possible side effects.

In a preferred embodiment, prostate cancer sequences are those that are upregulated in prostate cancer; that is, the expression of these genes is higher in the prostate cancer tissue as compared to non-cancerous tissue. "Up-regulation" as used herein often means at least about a two-fold change, preferably at least about a three fold change, with at least about five-fold or higher being preferred. All unigene cluster identification numbers and accession numbers herein are for the GenBank sequence database and the sequences of the accession numbers are hereby expressly incorporated by reference. GenBank is known in the art, see, e.g., Benson, DA, et al., Nucleic Acids Research 26:1-7 (1998) and http://www.ncbi.nlm.nih.gov/. Sequences are also available in other databases, e.g., European Molecular Biology Laboratory (EMBL) and DNA Database of Japan (DDBJ).

In another preferred embodiment, prostate cancer sequences are those that are down-regulated in prostate cancer; that is, the expression of these genes is lower in prostate

cancer tissue as compared to non-cancerous tissue (see, e.g., Tables 8, 12 and 14). "Down-regulation" as used herein often means at least about a 1.5-fold change more preferrably a two-fold change, preferably at least about a three fold change, with at least about five-fold or higher being most preferred.

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Informatics

The ability to identify genes that are over or under expressed in prostate cancer can additionally provide high-resolution, high-sensitivity datasets which can be used in the areas of diagnostics, therapeutics, drug development, pharmacogenetics, protein structure, biosensor development, and other related areas. For example, the expression profiles can be used in diagnostic or prognostic evaluation of patients with prostate cancer. Or as another example, subcellular toxicological information can be generated to better direct drug structure and activity correlation (see Anderson, Pharmaceutical Proteomics: Targets, Mechanism, and Function, paper presented at the IBC Proteomics conference, Coronado, CA (June 11-12, 1998)). Subcellular toxicological information can also be utilized in a biological sensor device to predict the likely toxicological effect of chemical exposures and likely tolerable exposure thresholds (see U.S. Patent No. 5,811,231). Similar advantages accrue from datasets relevant to other biomolecules and bioactive agents (e.g., nucleic acids, saccharides, lipids, drugs, and the like).

Thus, in another embodiment, the present invention provides a database that includes at least one set of assay data. The data contained in the database is acquired, e.g., using array analysis either singly or in a library format. The database can be in substantially any form in which data can be maintained and transmitted, but is preferably an electronic database. The electronic database of the invention can be maintained on any electronic device allowing for the storage of and access to the database, such as a personal computer, but is preferably distributed on a wide area network, such as the World Wide Web.

The focus of the present section on databases that include peptide sequence data is for clarity of illustration only. It will be apparent to those of skill in the art that similar databases can be assembled for any assay data acquired using an assay of the invention.

The compositions and methods for identifying and/or quantitating the relative and/or absolute abundance of a variety of molecular and macromolecular species from a biological sample undergoing prostate cancer, i.e., the identification of prostate cancer-associated sequences described herein, provide an abundance of information, which can be correlated with pathological conditions, predisposition to disease, drug testing, therapeutic monitoring, gene-disease causal linkages, identification of correlates of immunity and physiological status, among others. Although the data generated from the assays of the invention is suited for manual review and analysis, in a preferred embodiment, prior data processing using high-speed computers is utilized.

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An array of methods for indexing and retrieving biomolecular information is known in the art. For example, U.S. Patents 6,023,659 and 5,966,712 disclose a relational database system for storing biomolecular sequence information in a manner that allows sequences to be catalogued and searched according to one or more protein function hierarchies. U.S. Patent 5,953,727 discloses a relational database having sequence records containing information in a format that allows a collection of partial-length DNA sequences to be catalogued and searched according to association with one or more sequencing projects for obtaining full-length sequences from the collection of partial length sequences. U.S. Patent 5,706,498 discloses a gene database retrieval system for making a retrieval of a gene sequence similar to a sequence data item in a gene database based on the degree of similarity between a key sequence and a target sequence. U.S. Patent 5,538,897 discloses a method using mass spectroscopy fragmentation patterns of peptides to identify amino acid sequences in computer databases by comparison of predicted mass spectra with experimentally-derived mass spectra using a closeness-of-fit measure. U.S. Patent 5,926,818 discloses a multidimensional database comprising a functionality for multi-dimensional data analysis described as on-line analytical processing (OLAP), which entails the consolidation of projected and actual data according to more than one consolidation path or dimension. U.S. Patent 5,295,261 reports a hybrid database structure in which the fields of each database record are divided into two classes, navigational and informational data, with navigational fields stored in a hierarchical topological map which can be viewed as a tree structure or as the merger of two or more such tree structures.

See also Mount et al., Bioinformatics (2001); Biological Sequence Analysis: Probabilistic Models of Proteins and Nucleic Acids (Durbin et al., eds., 1999); Bioinformatics: A Practical Guide to the Analysis of Genes and Proteins (Baxevanis & Oeullette eds., 1998)); Rashidi & Buehler, Bioinformatics: Basic Applications in Biological Science and Medicine (1999); Introduction to Computational Molecular Biology (Setubal et al., eds 1997); Bioinformatics: Methods and Protocols (Misener & Krawetz, eds, 2000); Bioinformatics: Sequence, Structure, and Databanks: A Practical Approach (Higgins & Taylor, eds., 2000); Brown, Bioinformatics: A Biologist's Guide to Biocomputing and the Internet (2001); Han & Kamber, Data Mining: Concepts and Techniques (2000); and Waterman, Introduction to Computational Biology: Maps, Sequences, and Genomes (1995).

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The present invention provides a computer database comprising a computer and software for storing in computer-retrievable form assay data records cross-tabulated, e.g., with data specifying the source of the target-containing sample from which each sequence specificity record was obtained.

In an exemplary embodiment, at least one of the sources of target-containing sample is from a control tissue sample known to be free of pathological disorders. In a variation, at least one of the sources is a known pathological tissue specimen, e.g., a neoplastic lesion or another tissue specimen to be analyzed for prostate cancer. In another variation, the assay records cross-tabulate one or more of the following parameters for each target species in a sample: (1) a unique identification code, which can include, e.g., a target molecular structure and/or characteristic separation coordinate (e.g., electrophoretic coordinates); (2) sample source; and (3) absolute and/or relative quantity of the target species present in the sample.

The invention also provides for the storage and retrieval of a collection of target data in a computer data storage apparatus, which can include magnetic disks, optical disks, magneto-optical disks, DRAM, SRAM, SGRAM, SDRAM, RDRAM, DDR RAM, magnetic bubble memory devices, and other data storage devices, including CPU registers and on-CPU data storage arrays. Typically, the target data records are stored as a bit pattern in an array of magnetic domains on a magnetizable medium or as an array of charge states or transistor gate states, such as an array of cells in a DRAM device (e.g., each cell comprised of a transistor and a charge storage area, which may be on the transistor). In one embodiment,

the invention provides such storage devices, and computer systems built therewith, comprising a bit pattern encoding a protein expression fingerprint record comprising unique identifiers for at least 10 target data records cross-tabulated with target source.

When the target is a peptide or nucleic acid, the invention preferably provides a method for identifying related peptide or nucleic acid sequences, comprising performing a computerized comparison between a peptide or nucleic acid sequence assay record stored in or retrieved from a computer storage device or database and at least one other sequence. The comparison can include a sequence analysis or comparison algorithm or computer program embodiment thereof (e.g., FASTA, TFASTA, GAP, BESTFIT) and/or the comparison may be of the relative amount of a peptide or nucleic acid sequence in a pool of sequences determined from a polypeptide or nucleic acid sample of a specimen.

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The invention also preferably provides a magnetic disk, such as an IBM-compatible (DOS, Windows, Windows95/98/2000, Windows NT, OS/2) or other format (e.g., Linux, SunOS, Solaris, AIX, SCO Unix, VMS, MV, Macintosh, etc.) floppy diskette or hard (fixed, Winchester) disk drive, comprising a bit pattern encoding data from an assay of the invention in a file format suitable for retrieval and processing in a computerized sequence analysis, comparison, or relative quantitation method.

The invention also provides a network, comprising a plurality of computing devices linked via a data link, such as an Ethernet cable (coax or 10BaseT), telephone line, ISDN line, wireless network, optical fiber, or other suitable signal transmission medium, whereby at least one network device (e.g., computer, disk array, etc.) comprises a pattern of magnetic domains (e.g., magnetic disk) and/or charge domains (e.g., an array of DRAM cells) composing a bit pattern encoding data acquired from an assay of the invention.

The invention also provides a method for transmitting assay data that includes generating an electronic signal on an electronic communications device, such as a modem, ISDN terminal adapter, DSL, cable modem, ATM switch, or the like, wherein the signal includes (in native or encrypted format) a bit pattern encoding data from an assay or a database comprising a plurality of assay results obtained by the method of the invention.

In a preferred embodiment, the invention provides a computer system for comparing a query target to a database containing an array of data structures, such as an assay result obtained by the method of the invention, and ranking database targets based on the

degree of identity and gap weight to the target data. A central processor is preferably initialized to load and execute the computer program for alignment and/or comparison of the assay results. Data for a query target is entered into the central processor via an I/O device. Execution of the computer program results in the central processor retrieving the assay data from the data file, which comprises a binary description of an assay result.

The target data or record and the computer program can be transferred to secondary memory, which is typically random access memory (e.g., DRAM, SRAM, SGRAM, or SDRAM). Targets are ranked according to the degree of correspondence between a selected assay characteristic (e.g., binding to a selected affinity moiety) and the same characteristic of the query target and results are output via an I/O device. For example, a central processor can be a conventional computer (e.g., Intel Pentium, PowerPC, Alpha, PA-8000, SPARC, MIPS 4400, MIPS 10000, VAX, etc.); a program can be a commercial or public domain molecular biology software package (e.g., UWGCG Sequence Analysis Software, Darwin); a data file can be an optical or magnetic disk, a data server, a memory device (e.g., DRAM, SRAM, SGRAM, SDRAM, EPROM, bubble memory, flash memory, etc.); an I/O device can be a terminal comprising a video display and a keyboard, a modem, an ISDN terminal adapter, an Ethernet port, a punched card reader, a magnetic strip reader, or other suitable I/O device.

The invention also preferably provides the use of a computer system, such as that described above, which comprises: (1) a computer; (2) a stored bit pattern encoding a collection of peptide sequence specificity records obtained by the methods of the invention, which may be stored in the computer; (3) a comparison target, such as a query target; and (4) a program for alignment and comparison, typically with rank-ordering of comparison results on the basis of computed similarity values.

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Characteristics of prostate cancer-associated proteins

Prostate cancer proteins of the present invention may be classified as secreted proteins, transmembrane proteins or intracellular proteins. In one embodiment, the prostate cancer protein is an intracellular protein. Intracellular proteins may be found in the cytoplasm and/or in the nucleus. Intracellular proteins are involved in all aspects of cellular function and replication (including, e.g., signaling pathways); aberrant expression of such

proteins often results in unregulated or disregulated cellular processes (see, e.g., Molecular Biology of the Cell (Alberts, ed., 3rd ed., 1994). For example, many intracellular proteins have enzymatic activity such as protein kinase activity, protein phosphatase activity, protease activity, nucleotide cyclase activity, polymerase activity and the like. Intracellular proteins also serve as docking proteins that are involved in organizing complexes of proteins, or targeting proteins to various subcellular localizations, and are involved in maintaining the structural integrity of organelles.

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An increasingly appreciated concept in characterizing proteins is the presence in the proteins of one or more motifs for which defined functions have been attributed. In addition to the highly conserved sequences found in the enzymatic domain of proteins, highly conserved sequences have been identified in proteins that are involved in protein-protein interaction. For example, Src-homology-2 (SH2) domains bind tyrosine-phosphorylated targets in a sequence dependent manner. PTB domains, which are distinct from SH2 domains, also bind tyrosine phosphorylated targets. SH3 domains bind to proline-rich targets. In addition, PH domains, tetratricopeptide repeats and WD domains to name only a few, have been shown to mediate protein-protein interactions. Some of these may also be involved in binding to phospholipids or other second messengers. As will be appreciated by one of ordinary skill in the art, these motifs can be identified on the basis of primary sequence; thus, an analysis of the sequence of proteins may provide insight into both the enzymatic potential of the molecule and/or molecules with which the protein may associate. One useful database is Pfam (protein families), which is a large collection of multiple sequence alignments and hidden Markov models covering many common protein domains. Versions are available via the internet from Washington University in St. Louis, the Sanger Center in England, and the Karolinska Institute in Sweden (see, e.g., Bateman et al., Nuc. Acids Res. 28:263-266 (2000); Sonnhammer et al., Proteins 28:405-420 (1997); Bateman et al., Nuc. Acids Res. 27:260-262 (1999); and Sonnhammer et al., Nuc. Acids Res. 26:320-322-(1998)).

In another embodiment, the prostate cancer sequences are transmembrane proteins. Transmembrane proteins are molecules that span a phospholipid bilayer of a cell. They may have an intracellular domain, an extracellular domain, or both. The intracellular domains of such proteins may have a number of functions including those already described

for intracellular proteins. For example, the intracellular domain may have enzymatic activity and/or may serve as a binding site for additional proteins. Frequently the intracellular domain of transmembrane proteins serves both roles. For example certain receptor tyrosine kinases have both protein kinase activity and SH2 domains. In addition, autophosphorylation of tyrosines on the receptor molecule itself, creates binding sites for additional SH2 domain containing proteins.

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Transmembrane proteins may contain from one to many transmembrane domains. For example, receptor tyrosine kinases, certain cytokine receptors, receptor guanylyl cyclases and receptor serine/threonine protein kinases contain a single transmembrane domain. However, various other proteins including channels and adenylyl cyclases contain numerous transmembrane domains. Many important cell surface receptors such as G protein coupled receptors (GPCRs) are classified as "seven transmembrane domain" proteins, as they contain 7 membrane spanning regions. Characteristics of transmembrane domains include approximately 20 consecutive hydrophobic amino acids that may be followed by charged amino acids. Therefore, upon analysis of the amino acid sequence of a particular protein, the localization and number of transmembrane domains within the protein may be predicted (see, e.g. PSORT web site http://psort.nibb.ac.jp/). Important transmembrane protein receptors include, but are not limited to the insulin receptor, insulin-like growth factor receptor, human growth hormone receptor, glucose transporters, transferrin receptor, epidermal growth factor receptor, low density lipoprotein receptor, epidermal growth factor receptor, leptin receptor, interleukin receptors, e.g. IL-1 receptor, IL-2 receptor,

The extracellular domains of transmembrane proteins are diverse; however, conserved motifs are found repeatedly among various extracellular domains. Conserved structure and/or functions have been ascribed to different extracellular motifs. Many extracellular domains are involved in binding to other molecules. In one aspect, extracellular domains are found on receptors. Factors that bind the receptor domain include circulating ligands, which may be peptides, proteins, or small molecules such as adenosine and the like. For example, growth factors such as EGF, FGF and PDGF are circulating growth factors that bind to their cognate receptors to initiate a variety of cellular responses. Other factors include cytokines, mitogenic factors, neurotrophic factors and the like. Extracellular domains also

bind to cell-associated molecules. In this respect, they mediate cell-cell interactions. Cell-associated ligands can be tethered to the cell, e.g., via a glycosylphosphatidylinositol (GPI) anchor, or may themselves be transmembrane proteins. Extracellular domains also associate with the extracellular matrix and contribute to the maintenance of the cell structure.

Prostate cancer proteins that are transmembrane are particularly preferred in the present invention as they are readily accessible targets for immunotherapeutics, as are described herein. In addition, as outlined below, transmembrane proteins can be also useful in imaging modalities. Antibodies may be used to label such readily accessible proteins in situ. Alternatively, antibodies can also label intracellular proteins, in which case samples are typically permeablized to provide access to intracellular proteins.

It will also be appreciated by those in the art that a transmembrane protein can be made soluble by removing transmembrane sequences, e.g., through recombinant methods. Furthermore, transmembrane proteins that have been made soluble can be made to be secreted through recombinant means by adding an appropriate signal sequence.

In another embodiment, the prostate cancer proteins are secreted proteins; the secretion of which can be either constitutive or regulated. These proteins have a signal peptide or signal sequence that targets the molecule to the secretory pathway. Secreted proteins are involved in numerous physiological events; by virtue of their circulating nature, they serve to transmit signals to various other cell types. The secreted protein may function in an autocrine manner (acting on the cell that secreted the factor), a paracrine manner (acting on cells in close proximity to the cell that secreted the factor) or an endocrine manner (acting on cells at a distance). Thus secreted molecules find use in modulating or altering numerous aspects of physiology. Prostate cancer proteins that are secreted proteins are particularly preferred in the present invention as they serve as good targets for diagnostic markers, e.g., for blood, plasma, serum, or stool tests.

Use of prostate cancer nucleic acids

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As described above, prostate cancer sequence is initially identified by substantial nucleic acid and/or amino acid sequence homology or linkage to the prostate cancer sequences outlined herein. Such homology can be based upon the overall nucleic acid or amino acid sequence, and is generally determined as outlined below, using either

homology programs or hybridization conditions. Typically, linked sequences on a mRNA are found on the same molecule.

The prostate cancer nucleic acid sequences of the invention, e.g., the sequences in Tables 1-16, can be fragments of larger genes, i.e., they are nucleic acid segments. "Genes" in this context includes coding regions, non-coding regions, and mixtures of coding and non-coding regions. Accordingly, as will be appreciated by those in the art, using the sequences provided herein, extended sequences, in either direction, of the prostate cancer genes can be obtained, using techniques well known in the art for cloning either longer sequences or the full length sequences; see Ausubel, *et al.*, *supra*. Much can be done by informatics and many sequences can be clustered to include multiple sequences corresponding to a single gene, e.g., systems such as UniGene (see, http://www.ncbi.nlm.nih.gov/UniGene/).

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Once the prostate cancer nucleic acid is identified, it can be cloned and, if necessary, its constituent parts recombined to form the entire prostate cancer nucleic acid coding regions or the entire mRNA sequence. Once isolated from its natural source, e.g., contained within a plasmid or other vector or excised therefrom as a linear nucleic acid segment, the recombinant prostate cancer nucleic acid can be further-used as a probe to identify and isolate other prostate cancer nucleic acids, e.g., extended coding regions. It can also be used as a "precursor" nucleic acid to make modified or variant prostate cancer nucleic acids and proteins.

The prostate cancer nucleic acids of the present invention are used in several ways. In a first embodiment, nucleic acid probes to the prostate cancer nucleic acids are made and attached to biochips to be used in screening and diagnostic methods, as outlined below, or for administration, e.g., for gene therapy, vaccine, and/or antisense applications. Alternatively, the prostate cancer nucleic acids that include coding regions of prostate cancer proteins can be put into expression vectors for the expression of prostate cancer proteins, again for screening purposes or for administration to a patient.

In a preferred embodiment, nucleic acid probes to prostate cancer nucleic acids (both the nucleic acid sequences outlined in the figures and/or the complements thereof) are made. The nucleic acid probes attached to the biochip are designed to be substantially complementary to the prostate cancer nucleic acids, *i.e.* the target sequence (either the target

sequence of the sample or to other probe sequences, e.g., in sandwich assays), such that hybridization of the target sequence and the probes of the present invention occurs. As outlined below, this complementarity need not be perfect; there may be any number of base pair mismatches which will interfere with hybridization between the target sequence and the single stranded nucleic acids of the present invention. However, if the number of mutations is so great that no hybridization can occur under even the least stringent of hybridization conditions, the sequence is not a complementary target sequence. Thus, by "substantially complementary" herein is meant that the probes are sufficiently complementary to the target sequences to hybridize under normal reaction conditions, particularly high stringency conditions, as outlined herein.

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A nucleic acid probe is generally single stranded but can be partially single and partially double stranded. The strandedness of the probe is dictated by the structure, composition, and properties of the target sequence. In general, the nucleic acid probes range from about 8 to about 100 bases long, with from about 10 to about 80 bases being preferred, and from about 30 to about 50 bases being particularly preferred. That is, generally whole genes are not used. In some embodiments, much longer nucleic acids can be used, up to hundreds of bases.

In a preferred embodiment, more than one probe per sequence is used, with either overlapping probes or probes to different sections of the target being used. That is, two, three, four or more probes, with three being preferred, are used to build in a redundancy for a particular target. The probes can be overlapping (i.e., have some sequence in common), or separate. In some cases, PCR primers may be used to amplify signal for higher sensitivity.

As will be appreciated by those in the art, nucleic acids can be attached or immobilized to a solid support in a wide variety of ways. By "immobilized" and grammatical equivalents herein is meant the association or binding between the nucleic acid probe and the solid support is sufficient to be stable under the conditions of binding, washing, analysis, and removal as outlined below. The binding can typically be covalent or non-covalent. By "non-covalent binding" and grammatical equivalents herein is meant one or more of electrostatic, hydrophilic, and hydrophobic interactions. Included in non-covalent binding is the covalent attachment of a molecule, such as, streptavidin to the support and the non-covalent binding of the biotinylated probe to the streptavidin. By "covalent binding" and grammatical

equivalents herein is meant that the two moieties, the solid support and the probe, are attached by at least one bond, including sigma bonds, pi bonds and coordination bonds. Covalent bonds can be formed directly between the probe and the solid support or can be formed by a cross linker or by inclusion of a specific reactive group on either the solid support or the probe or both molecules. Immobilization may also involve a combination of covalent and non-covalent interactions.

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In general, the probes are attached to the biochip in a wide variety of ways, as will be appreciated by those in the art. As described herein, the nucleic acids can either be synthesized first, with subsequent attachment to the biochip, or can be directly synthesized on the biochip.

The biochip comprises a suitable solid substrate. By "substrate" or "solid support" or other grammatical equivalents herein is meant a material that can be modified to contain discrete individual sites appropriate for the attachment or association of the nucleic acid probes and is amenable to at least one detection method. As will be appreciated by those in the art, the number of possible substrates are very large, and include, but are not limited to, glass and modified or functionalized glass, plastics (including acrylics, polystyrene and copolymers of styrene and other materials, polypropylene, polyethylene, polybutylene, polyurethanes, TeflonJ, etc.), polysaccharides, nylon or nitrocellulose, resins, silica or silicabased materials including silicon and modified silicon, carbon, metals, inorganic glasses, plastics, etc. In general, the substrates allow optical detection and do not appreciably fluoresce. A preferred substrate is described in copending application entitled Reusable Low Fluorescent Plastic Biochip, U.S. Application Serial No. 09/270,214, filed March 15, 1999, herein incorporated by reference in its entirety.

Generally the substrate is planar, although as will be appreciated by those in the art, other configurations of substrates may be used as well. For example, the probes may be placed on the inside surface of a tube, for flow-through sample analysis to minimize sample volume. Similarly, the substrate may be flexible, such as a flexible foam, including closed cell foams made of particular plastics.

In a preferred embodiment, the surface of the biochip and the probe may be derivatized with chemical functional groups for subsequent attachment of the two. Thus, e.g., the biochip is derivatized with a chemical functional group including, but not limited to,

amino groups, carboxy groups, oxo groups and thiol groups, with amino groups being particularly preferred. Using these functional groups, the probes can be attached using functional groups on the probes. For example, nucleic acids containing amino groups can be attached to surfaces comprising amino groups, e.g. using linkers as are known in the art; e.g., homo-or hetero-bifunctional linkers as are well known (see 1994 Pierce Chemical Company catalog, technical section on cross-linkers, pages 155-200). In addition, in some cases, additional linkers, such as alkyl groups (including substituted and heteroalkyl groups) may be used.

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In this embodiment, oligonucleotides are synthesized as is known in the art, and then attached to the surface of the solid support. As will be appreciated by those skilled in the art, either the 5' or 3' terminus may be attached to the solid support, or attachment may be via an internal nucleoside.

In another embodiment, the immobilization to the solid support may be very strong, yet non-covalent. For example, biotinylated oligonucleotides can be made, which bind to surfaces covalently coated with streptavidin, resulting in attachment.

Alternatively, the oligonucleotides may be synthesized on the surface, as is known in the art. For example, photoactivation techniques utilizing photopolymerization compounds and techniques are used. In a preferred embodiment, the nucleic acids can be synthesized in situ, using well known photolithographic techniques, such as those described in WO 95/25116; WO 95/35505; U.S. Patent Nos. 5,700,637 and 5,445,934; and references cited within, all of which are expressly incorporated by reference; these methods of attachment form the basis of the Affimetrix GeneChipTM technology.

Often, amplification-based assays are performed to measure the expression level of prostate cancer-associated sequences. These assays are typically performed in conjunction with reverse transcription. In such assays, a prostate cancer-associated nucleic acid sequence acts as a template in an amplification reaction (e.g., Polymerase Chain Reaction, or PCR). In a quantitative amplification, the amount of amplification product will be proportional to the amount of template in the original sample. Comparison to appropriate controls provides a measure of the amount of prostate cancer-associated RNA. Methods of quantitative amplification are well known to those of skill in the art. Detailed protocols for

quantitative PCR are provided, e.g., in Innis et al., PCR Protocols, A Guide to Methods and Applications (1990).

In some embodiments, a TaqMan based assay is used to measure expression. TaqMan based assays use a fluorogenic oligonucleotide probe that contains a 5' fluorescent dye and a 3' quenching agent. The probe hybridizes to a PCR product, but cannot itself be extended due to a blocking agent at the 3' end. When the PCR product is amplified in subsequent cycles, the 5' nuclease activity of the polymerase, e.g., AmpliTaq, results in the cleavage of the TaqMan probe. This cleavage separates the 5' fluorescent dye and the 3' quenching agent, thereby resulting in an increase in fluorescence as a function of amplification (see, e.g., literature provided by Perkin-Elmer, e.g., www2.perkin-elmer.com).

Other suitable amplification methods include, but are not limited to, ligase chain reaction (LCR) (see Wu & Wallace, Genomics 4:560 (1989), Landegren et al., Science 241:1077 (1988), and Barringer et al., Gene 89:117 (1990)), transcription amplification (Kwoh et al., Proc. Natl. Acad. Sci. USA 86:1173 (1989)), self-sustained sequence replication (Guatelli et al., Proc. Nat. Acad. Sci. USA 87:1874 (1990)), dot PCR, and linker adapter PCR, etc.

Expression of prostate cancer proteins from nucleic acids

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In a preferred embodiment, prostate cancer nucleic acids, e.g., encoding prostate cancer proteins are used to make a variety of expression vectors to express prostate cancer proteins which can then be used in screening assays, as described below. Expression vectors and recombinant DNA technology are well known to those of skill in the art (see, e.g., Ausubel, supra, and Gene Expression Systems (Fernandez & Hoeffler, eds, 1999)) and are used to express proteins. The expression vectors may be either self-replicating extrachromosomal vectors or vectors which integrate into a host genome. Generally, these expression vectors include transcriptional and translational regulatory nucleic acid operably linked to the nucleic acid encoding the prostate cancer protein. The term "control sequences" refers to DNA sequences used for the expression of an operably linked coding sequence in a particular host organism. Control sequences that are suitable for prokaryotes, e.g., include a promoter, optionally an operator sequence, and a ribosome binding site. Eukaryotic cells are known to utilize promoters, polyadenylation signals, and enhancers.

Nucleic acid is "operably linked" when it is placed into a functional relationship with another nucleic acid sequence. For example, DNA for a presequence or secretory leader is operably linked to DNA for a polypeptide if it is expressed as a preprotein that participates in the secretion of the polypeptide; a promoter or enhancer is operably linked to a coding sequence if it affects the transcription of the sequence; or a ribosome binding site is operably linked to a coding sequence if it is positioned so as to facilitate translation.

Generally, "operably linked" means that the DNA sequences being linked are contiguous, and, in the case of a secretory leader, contiguous and in reading phase. However, enhancers do not have to be contiguous. Linking is typically accomplished by ligation at convenient restriction sites. If such sites do not exist, synthetic oligonucleotide adaptors or linkers are used in accordance with conventional practice. Transcriptional and translational regulatory nucleic acid will generally be appropriate to the host cell used to express the prostate cancer protein. Numerous types of appropriate expression vectors, and suitable regulatory sequences are known in the art for a variety of host cells.

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In general, transcriptional and translational regulatory sequences may include, but are not limited to, promoter sequences, ribosomal binding sites, transcriptional start and stop sequences, translational start and stop sequences, and enhancer or activator sequences. In a preferred embodiment, the regulatory sequences include a promoter and transcriptional start and stop sequences.

Promoter sequences encode either constitutive or inducible promoters. The promoters may be either naturally occurring promoters or hybrid promoters. Hybrid promoters, which combine elements of more than one promoter, are also known in the art, and are useful in the present invention.

In addition, an expression vector may comprise additional elements. For example, the expression vector may have two replication systems, thus allowing it to be maintained in two organisms, e.g. in mammalian or insect cells for expression and in a procaryotic host for cloning and amplification. Furthermore, for integrating expression vectors, the expression vector contains at least one sequence homologous to the host cell genome, and preferably two homologous sequences which flank the expression construct. The integrating vector may be directed to a specific locus in the host cell by selecting the

appropriate homologous sequence for inclusion in the vector. Constructs for integrating vectors are well known in the art (e.g., Fernandez & Hoeffler, *supra*).

In addition, in a preferred embodiment, the expression vector contains a selectable marker gene to allow the selection of transformed host cells. Selection genes are well known in the art and will vary with the host cell used.

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The prostate cancer proteins of the present invention are produced by culturing a host cell transformed with an expression vector containing nucleic acid encoding a prostate cancer protein, under the appropriate conditions to induce or cause expression of the prostate cancer protein. Conditions appropriate for prostate cancer protein expression will vary with the choice of the expression vector and the host cell, and will be easily ascertained by one skilled in the art through routine experimentation or optimization. For example, the use of constitutive promoters in the expression vector will require optimizing the growth and proliferation of the host cell, while the use of an inducible promoter requires the appropriate growth conditions for induction. In addition, in some embodiments, the timing of the harvest is important. For example, the baculoviral systems used in insect cell expression are lytic viruses, and thus harvest time selection can be crucial for product yield.

Appropriate host cells include yeast, bacteria, archaebacteria, fungi, and insect and animal cells, including mammalian cells. Of particular interest are *Saccharomyces* cerevisiae and other yeasts, *E. coli, Bacillus subtilis*, Sf9 cells, C129 cells, 293 cells, *Neurospora*, BHK, CHO, COS, HeLa cells, HUVEC (human umbilical vein endothelial cells), THP1 cells (a macrophage cell line) and various other human cells and cell lines.

In a preferred embodiment, the prostate cancer proteins are expressed in mammalian cells. Mammalian expression systems are also known in the art, and include retroviral and adenoviral systems. One expression vector system is a retroviral vector system such as is generally described in PCT/US97/01019 and PCT/US97/01048, both of which are hereby expressly incorporated by reference. Of particular use as mammalian promoters are the promoters from mammalian viral genes, since the viral genes are often highly expressed and have a broad host range. Examples include the SV40 early promoter, mouse mammary tumor virus LTR promoter, adenovirus major late promoter, herpes simplex virus promoter, and the CMV promoter (see, e.g., Fernandez & Hoeffler, supra). Typically, transcription termination and polyadenylation sequences recognized by mammalian cells are regulatory

regions located 3' to the translation stop codon and thus, together with the promoter elements, flank the coding sequence. Examples of transcription terminator and polyadenlyation signals include those derived form SV40.

The methods of introducing exogenous nucleic acid into mammalian hosts, as well as other hosts, is well known in the art, and will vary with the host cell used.

Techniques include dextran-mediated transfection, calcium phosphate precipitation, polybrene mediated transfection, protoplast fusion, electroporation, viral infection, encapsulation of the polynucleotide(s) in liposomes, and direct microinjection of the DNA into nuclei.

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In a preferred embodiment, prostate cancer proteins are expressed in bacterial systems. Bacterial expression systems are well known in the art. Promoters from bacteriophage may also be used and are known in the art. In addition, synthetic promoters and hybrid promoters are also useful; e.g., the tac promoter is a hybrid of the trp and lac promoter sequences. Furthermore, a bacterial promoter can include naturally occurring promoters of non-bacterial origin that have the ability to bind bacterial RNA polymerase and initiate transcription. In addition to a functioning promoter sequence, an efficient ribosome binding site is desirable. The expression vector may also include a signal peptide sequence that provides for secretion of the prostate cancer protein in bacteria. The protein is either secreted into the growth media (gram-positive bacteria) or into the periplasmic space, located between the inner and outer membrane of the cell (gram-negative bacteria). The bacterial expression vector may also include a selectable marker gene to allow for the selection of bacterial strains that have been transformed. Suitable selection genes include genes which render the bacteria resistant to drugs such as ampicillin, chloramphenicol, erythromycin, kanamycin, neomycin and tetracycline. Selectable markers also include biosynthetic genes, such as those in the histidine, tryptophan and leucine biosynthetic pathways. These components are assembled into expression vectors. Expression vectors for bacteria are well known in the art, and include vectors for Bacillus subtilis, E. coli, Streptococcus cremoris, and Streptococcus lividans, among others (e.g., Fernandez & Hoeffler, supra). The bacterial expression vectors are transformed into bacterial host cells using techniques well known in the art, such as calcium chloride treatment, electroporation, and others.

In one embodiment, prostate cancer proteins are produced in insect cells.

Expression vectors for the transformation of insect cells, and in particular, baculovirus-based expression vectors, are well known in the art.

In a preferred embodiment, prostate cancer protein is produced in yeast cells.

Yeast expression systems are well known in the art, and include expression vectors for Saccharomyces cerevisiae, Candida albicans and C. maltosa, Hansenula polymorpha, Kluyveromyces fragilis and K. lactis, Pichia guillerimondii and P. pastoris, Schizosaccharomyces pombe, and Yarrowia lipolytica.

The prostate cancer protein may also be made as a fusion protein, using techniques well known in the art. Thus, e.g., for the creation of monoclonal antibodies, if the desired epitope is small, the prostate cancer protein may be fused to a carrier protein to form an immunogen. Alternatively, the prostate cancer protein may be made as a fusion protein to increase expression, or for other reasons. For example, when the prostate cancer protein is a prostate cancer peptide, the nucleic acid encoding the peptide may be linked to other nucleic acid for expression purposes.

In a preferred embodiment, the prostate cancer protein is purified or isolated after expression. Prostate cancer proteins may be isolated or purified in a variety of ways known to those skilled in the art depending on what other components are present in the sample. Standard purification methods include electrophoretic, molecular, immunological and chromatographic techniques, including ion exchange, hydrophobic, affinity, and reverse-phase HPLC chromatography, and chromatofocusing. For example, the prostate cancer protein may be purified using a standard anti-prostate cancer protein antibody column. Ultrafiltration and diafiltration techniques, in conjunction with protein concentration, are also useful. For general guidance in suitable purification techniques, see Scopes, *Protein Purification* (1982). The degree of purification necessary will vary depending on the use of the prostate cancer protein. In some instances no purification will be necessary.

Once expressed and purified if necessary, the prostate cancer proteins and nucleic acids are useful in a number of applications. They may be used as immunoselection reagents, as vaccine reagents, as screening agents, etc.

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Variants of prostate cancer proteins

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In one embodiment, the prostate cancer proteins are derivative or variant prostate cancer proteins as compared to the wild-type sequence. That is, as outlined more fully below, the derivative prostate cancer peptide will often contain at least one amino acid substitution, deletion or insertion, with amino acid substitutions being particularly preferred. The amino acid substitution, insertion or deletion may occur at any residue within the prostate cancer peptide.

Also included within one embodiment of prostate cancer proteins of the present invention are amino acid sequence variants. These variants typically fall into one or more of three classes: substitutional, insertional or deletional variants. These variants ordinarily are prepared by site specific mutagenesis of nucleotides in the DNA encoding the prostate cancer protein, using cassette or PCR mutagenesis or other techniques well known in the art, to produce DNA encoding the variant, and thereafter expressing the DNA in recombinant cell culture as outlined above. However, variant prostate cancer protein fragments having up to about 100-150 residues may be prepared by in vitro synthesis using established techniques. Amino acid sequence variants are characterized by the predetermined nature of the variation, a feature that sets them apart from naturally occurring allelic or interspecies variation of the prostate cancer protein amino acid sequence. The variants typically exhibit the same qualitative biological activity as the naturally occurring analogue, although variants can also be selected which have modified characteristics as will be more fully outlined below.

While the site or region for introducing an amino acid sequence variation is predetermined, the mutation per se need not be predetermined. For example, in order to optimize the performance of a mutation at a given site, random mutagenesis may be conducted at the target codon or region and the expressed prostate cancer variants screened for the optimal combination of desired activity. Techniques for making substitution mutations at predetermined sites in DNA having a known sequence are well known, e.g., M13 primer mutagenesis and PCR mutagenesis. Screening of the mutants is done using assays of prostate cancer protein activities.

Amino acid substitutions are typically of single residues; insertions usually will be on the order of from about 1 to 20 amino acids, although considerably larger

insertions may be tolerated. Deletions range from about 1 to about 20 residues, although in some cases deletions may be much larger.

Substitutions, deletions, insertions or any combination thereof may be used to arrive at a final derivative. Generally these changes are done on a few amino acids to minimize the alteration of the molecule. However, larger changes may be tolerated in certain circumstances. When small alterations in the characteristics of the prostate cancer protein are desired, substitutions are generally made in accordance with the amino acid substitution relationships provided in the definition section.

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The variants typically exhibit the same qualitative biological activity and will elicit the same immune response as the naturally-occurring analog, although variants also are selected to modify the characteristics of the prostate cancer proteins as needed. Alternatively, the variant may be designed such that the biological activity of the prostate cancer protein is altered. For example, glycosylation sites may be altered or removed.

Substantial changes in function or immunological identity are made by selecting substitutions that are less conservative than those described above. For example, substitutions may be made which more significantly affect: the structure of the polypeptide backbone in the area of the alteration, for example the alpha-helical or beta-sheet structure; the charge or hydrophobicity of the molecule at the target site; or the bulk of the side chain. The substitutions which in general are expected to produce the greatest changes in the polypeptide's properties are those in which (a) a hydrophilic residue, e.g. seryl or threonyl is substituted for (or by) a hydrophobic residue, e.g. leucyl, isoleucyl, phenylalanyl, valyl or alanyl; (b) a cysteine or proline is substituted for (or by) any other residue; (c) a residue having an electropositive side chain, e.g. lysyl, arginyl, or histidyl, is substituted for (or by) an electronegative residue, e.g. glutamyl or aspartyl; or (d) a residue having a bulky side chain, e.g. phenylalanine, is substituted for (or by) one not having a side chain, e.g. glycine.

Covalent modifications of prostate cancer polypeptides are included within the scope of this invention. One type of covalent modification includes reacting targeted amino acid residues of a prostate cancer polypeptide with an organic derivatizing agent that is capable of reacting with selected side chains or the N-or C-terminal residues of a prostate cancer polypeptide. Derivatization with bifunctional agents is useful, for instance, for crosslinking prostate cancer polypeptides to a water-insoluble support matrix or surface for

use in the method for purifying anti-prostate cancer polypeptide antibodies or screening assays, as is more fully described below. Commonly used crosslinking agents include, e.g., 1,1-bis(diazoacetyl)-2-phenylethane, glutaraldehyde, N-hydroxysuccinimide esters, e.g., esters with 4-azidosalicylic acid, homobifunctional imidoesters, including disuccinimidyl esters such as 3,3'-dithiobis(succinimidylpropionate), bifunctional maleimides such as bis-N-maleimido-1,8-octane and agents such as methyl-3-((p-azidophenyl)dithio)propioimidate.

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Other modifications include deamidation of glutaminyl and asparaginyl residues to the corresponding glutamyl and aspartyl residues, respectively, hydroxylation of proline and lysine, phosphorylation of hydroxyl groups of seryl, threonyl or tyrosyl residues, methylation of the amino groups of the lysine, arginine, and histidine side chains (Creighton, *Proteins: Structure and Molecular Properties*, pp. 79-86 (1983)), acetylation of the N-terminal amine, and amidation of any C-terminal carboxyl group.

Another type of covalent modification of the prostate cancer polypeptide included within the scope of this invention comprises altering the native glycosylation pattern of the polypeptide. "Altering the native glycosylation pattern" is intended for purposes herein to mean deleting one or more carbohydrate moieties found in native sequence prostate cancer polypeptide, and/or adding one or more glycosylation sites that are not present in the native sequence prostate cancer polypeptide. Glycosylation patterns can be altered in many ways. For example the use of different cell types to express prostate cancer-associated sequences can result in different glycosylation patterns.

Addition of glycosylation sites to prostate cancer polypeptides may also be accomplished by altering the amino acid sequence thereof. The alteration may be made, e.g., by the addition of, or substitution by, one or more serine or threonine residues to the native sequence prostate cancer polypeptide (for O-linked glycosylation sites). The prostate cancer amino acid sequence may optionally be altered through changes at the DNA level, particularly by mutating the DNA encoding the prostate cancer polypeptide at preselected bases such that codons are generated that will translate into the desired amino acids.

Another means of increasing the number of carbohydrate moieties on the prostate cancer polypeptide is by chemical or enzymatic coupling of glycosides to the polypeptide. Such methods are described in the art, e.g., in WO 87/05330, and in Aplin & Wriston, CRC Crit. Rev. Biochem., pp. 259-306 (1981).

Removal of carbohydrate moieties present on the prostate cancer polypeptide may be accomplished chemically or enzymatically or by mutational substitution of codons encoding for amino acid residues that serve as targets for glycosylation. Chemical deglycosylation techniques are known in the art and described, for instance, by Hakimuddin, et al., Arch. Biochem. Biophys., 259:52 (1987) and by Edge et al., Anal. Biochem., 118:131 (1981). Enzymatic cleavage of carbohydrate moieties on polypeptides can be achieved by the use of a variety of endo-and exo-glycosidases as described by Thotakura et al., Meth. Enzymol., 138:350 (1987).

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Another type of covalent modification of prostate cancer comprises linking the prostate cancer polypeptide to one of a variety of nonproteinaceous polymers, e.g., polyethylene glycol, polypropylene glycol, or polyoxyalkylenes, in the manner set forth in U.S. Patent Nos. 4,640,835; 4,496,689; 4,301,144; 4,670,417; 4,791,192 or 4,179,337.

Prostate cancer polypeptides of the present invention may also be modified in a way to form chimeric molecules comprising a prostate cancer polypeptide fused to another, heterologous polypeptide or amino acid sequence. In one embodiment, such a chimeric molecule comprises a fusion of a prostate cancer polypeptide with a tag polypeptide which provides an epitope to which an anti-tag antibody can selectively bind. The epitope tag is generally placed at the amino-or carboxyl-terminus of the prostate cancer polypeptide. The presence of such epitope-tagged forms of a prostate cancer polypeptide can be detected using an antibody against the tag polypeptide. Also, provision of the epitope tag enables the prostate cancer polypeptide to be readily purified by affinity purification using an anti-tag antibody or another type of affinity matrix that binds to the epitope tag. In an alternative embodiment, the chimeric molecule may comprise a fusion of a prostate cancer polypeptide with an immunoglobulin or a particular region of an immunoglobulin. For a bivalent form of the chimeric molecule, such a fusion could be to the Fc region of an IgG molecule.

Various tag polypeptides and their respective antibodies are well known in the art. Examples include poly-histidine (poly-his) or poly-histidine-glycine (poly-his-gly) tags; HIS6 and metal chelation tags, the flu HA tag polypeptide and its antibody 12CA5 (Field et al., Mol. Cell. Biol. 8:2159-2165 (1988)); the c-myc tag and the 8F9, 3C7, 6E10, G4, B7 and 9E10 antibodies thereto (Evan et al., Molecular and Cellular Biology 5:3610-3616 (1985)); and the Herpes Simplex virus glycoprotein D (gD) tag and its antibody (Paborsky et al.,

Protein Engineering 3(6):547-553 (1990)). Other tag polypeptides include the Flag-peptide (Hopp et al., BioTechnology 6:1204-1210 (1988)); the KT3 epitope peptide (Martin et al., Science 255:192-194 (1992)); tubulin epitope peptide (Skinner et al., J. Biol. Chem. 266:15163-15166 (1991)); and the T7 gene 10 protein peptide tag (Lutz-Freyermuth et al., Proc. Natl. Acad. Sci. USA 87:6393-6397 (1990)).

Also included are other prostate cancer proteins of the prostate cancer family, and prostate cancer proteins from other organisms, which are cloned and expressed as outlined below. Thus, probe or degenerate polymerase chain reaction (PCR) primer sequences may be used to find other related prostate cancer proteins from humans or other organisms. As will be appreciated by those in the art, particularly useful probe and/or PCR primer sequences include the unique areas of the prostate cancer nucleic acid sequence. As is generally known in the art, preferred PCR primers are from about 15 to about 35 nucleotides in length, with from about 20 to about 30 being preferred, and may contain inosine as needed. The conditions for the PCR reaction are well known in the art (e.g., Innis, PCR Protocols, supra).

Antibodies to prostate cancer proteins

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In a preferred embodiment, when the prostate cancer protein is to be used to generate antibodies, e.g., for immunotherapy or immunodiagnosis, the prostate cancer protein should share at least one epitope or determinant with the full length protein. By "epitope" or "determinant" herein is typically meant a portion of a protein which will generate and/or bind an antibody or T-cell receptor in the context of MHC. Thus, in most instances, antibodies made to a smaller prostate cancer protein will be able to bind to the full-length protein, particularly linear epitopes. In a preferred embodiment, the epitope is unique; that is, antibodies generated to a unique epitope show little or no cross-reactivity.

Methods of preparing polyclonal antibodies are known to the skilled artisan (e.g., Coligan, supra; and Harlow & Lane, supra). Polyclonal antibodies can be raised in a mammal, e.g., by one or more injections of an immunizing agent and, if desired, an adjuvant. Typically, the immunizing agent and/or adjuvant will be injected in the mammal by multiple subcutaneous or intraperitoneal injections. The immunizing agent may include a protein encoded by a nucleic acid of the figures or fragment thereof or a fusion protein thereof. It

may be useful to conjugate the immunizing agent to a protein known to be immunogenic in the mammal being immunized. Examples of such immunogenic proteins include but are not limited to keyhole limpet hemocyanin, serum albumin, bovine thyroglobulin, and soybean trypsin inhibitor. Examples of adjuvants which may be employed include Freund's complete adjuvant and MPL-TDM adjuvant (monophosphoryl Lipid A, synthetic trehalose dicorynomycolate). The immunization protocol may be selected by one skilled in the art without undue experimentation.

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The antibodies may, alternatively, be monoclonal antibodies. Monoclonal antibodies may be prepared using hybridoma methods, such as those described by Kohler & Milstein, Nature 256:495 (1975). In a hybridoma method, a mouse, hamster, or other appropriate host animal, is typically immunized with an immunizing agent to elicit lymphocytes that produce or are capable of producing antibodies that will specifically bind to the immunizing agent. Alternatively, the lymphocytes may be immunized in vitro. The immunizing agent will typically include a polypeptide encoded by a nucleic acid of Tables 1-16 fragment thereof, or a fusion protein thereof. Generally, either peripheral blood lymphocytes ("PBLs") are used if cells of human origin are desired, or spleen cells or lymph node cells are used if non-human mammalian sources are desired. The lymphocytes are then fused with an immortalized cell line using a suitable fusing agent, such as polyethylene glycol, to form a hybridoma cell (Goding, Monoclonal Antibodies: Principles and Practice, pp. 59-103 (1986)). Immortalized cell lines are usually transformed mammalian cells, particularly myeloma cells of rodent, bovine and human origin. Usually, rat or mouse myeloma cell lines are employed. The hybridoma cells may be cultured in a suitable culture medium that preferably contains one or more substances that inhibit the growth or survival of the unfused, immortalized cells. For example, if the parental cells lack the enzyme hypoxanthine guanine phosphoribosyl transferase (HGPRT or HPRT), the culture medium for the hybridomas typically will include hypoxanthine, aminopterin, and thymidine ("HAT medium"), which substances prevent the growth of HGPRT-deficient cells.

In one embodiment, the antibodies are bispecific antibodies. Bispecific antibodies are monoclonal, preferably human or humanized, antibodies that have binding specificities for at least two different antigens or that have binding specificities for two epitopes on the same antigen. In one embodiment, one of the binding specificities is for a

protein encoded by a nucleic acid Tables 1-16 or a fragment thereof, the other one is for any other antigen, and preferably for a cell-surface protein or receptor or receptor subunit, preferably one that is tumor specific. Alternatively, tetramer-type technology may create multivalent reagents.

In a preferred embodiment, the antibodies to prostate cancer protein are capable of reducing or eliminating a biological function of a prostate cancer protein, as is described below. That is, the addition of anti-prostate cancer protein antibodies (either polyclonal or preferably monoclonal) to prostate cancer tissue (or cells containing prostate cancer) may reduce or eliminate the prostate cancer. Generally, at least a 25% decrease in activity, growth, size or the like is preferred, with at least about 50% being particularly preferred and about a 95-100% decrease being especially preferred.

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In a preferred embodiment the antibodies to the prostate cancer proteins are humanized antibodies (e.g., Xenerex Biosciences, Mederex, Inc., Abgenix, Inc., Protein Design Labs, Inc.) Humanized forms of non-human (e.g., murine) antibodies are chimeric molecules of immunoglobulins, immunoglobulin chains or fragments thereof (such as Fv, Fab, Fab', F(ab')2 or other antigen-binding subsequences of antibodies) which contain minimal sequence derived from non-human immunoglobulin. Humanized antibodies include human immunoglobulins (recipient antibody) in which residues from a complementary determining region (CDR) of the recipient are replaced by residues from a CDR of a nonhuman species (donor antibody) such as mouse, rat or rabbit having the desired specificity, affinity and capacity. In some instances, Fv framework residues of the human immunoglobulin are replaced by corresponding non-human residues. Humanized antibodies may also comprise residues which are found neither in the recipient antibody nor in the imported CDR or framework sequences. In general, a humanized antibody will comprise substantially all of at least one, and typically two, variable domains, in which all or substantially all of the CDR regions correspond to those of a non-human immunoglobulin and all or substantially all of the framework (FR) regions are those of a human immunoglobulin consensus sequence. The humanized antibody optimally also will comprise at least a portion of an immunoglobulin constant region (Fc), typically that of a human immunoglobulin (Jones et al., Nature 321:522-525 (1986); Riechmann et al., Nature 332:323-329 (1988); and Presta, Curr. Op. Struct. Biol. 2:593-596 (1992)). Humanization

can be essentially performed following the method of Winter and co-workers (Jones et al., Nature 321:522-525 (1986); Riechmann et al., Nature 332:323-327 (1988); Verhoeyen et al., Science 239:1534-1536 (1988)), by substituting rodent CDRs or CDR sequences for the corresponding sequences of a human antibody. Accordingly, such humanized antibodies are chimeric antibodies (U.S. Patent No. 4,816,567), wherein substantially less than an intact human variable domain has been substituted by the corresponding sequence from a non-human species.

Human antibodies can also be produced using various techniques known in the art, including phage display libraries (Hoogenboom & Winter, *J. Mol. Biol.* 227:381 (1991); Marks et al., *J. Mol. Biol.* 222:581 (1991)). The techniques of Cole et al. and Boerner et al. are also available for the preparation of human monoclonal antibodies (Cole et al., *Monoclonal Antibodies and Cancer Therapy*, p. 77 (1985) and Boerner et al., *J. Immunol.* 147(1):86-95 (1991)). Similarly, human antibodies can be made by introducing of human immunoglobulin loci into transgenic animals, e.g., mice in which the endogenous immunoglobulin genes have been partially or completely inactivated. Upon challenge, human antibody production is observed, which closely resembles that seen in humans in all respects, including gene rearrangement, assembly, and antibody repertoire. This approach is described, e.g., in U.S. Patent Nos. 5,545,807; 5,545,806; 5,569,825; 5,625,126; 5,633,425; 5,661,016, and in the following scientific publications: Marks et al., Bio/Technology 10:779-783 (1992); Lonberg et al., Nature 368:856-859 (1994); Morrison, Nature 368:812-13 (1994); Fishwild et al., Nature Biotechnology 14:845-51 (1996); Neuberger, Nature Biotechnology 14:826 (1996); Lonberg & Huszar, Intern. Rev. Immunol. 13:65-93 (1995).

By immunotherapy is meant treatment of prostate cancer with an antibody raised against prostate cancer proteins. As used herein, immunotherapy can be passive or active. Passive immunotherapy as defined herein is the passive transfer of antibody to a recipient (patient). Active immunization is the induction of antibody and/or T-cell responses in a recipient (patient). Induction of an immune response is the result of providing the recipient with an antigen to which antibodies are raised. As appreciated by one of ordinary skill in the art, the antigen may be provided by injecting a polypeptide against which antibodies are desired to be raised into a recipient, or contacting the recipient with a nucleic

acid capable of expressing the antigen and under conditions for expression of the antigen, leading to an immune response.

In a preferred embodiment the prostate cancer proteins against which antibodies are raised are secreted proteins as described above. Without being bound by theory, antibodies used for treatment, bind and prevent the secreted protein from binding to its receptor, thereby inactivating the secreted prostate cancer protein.

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In another preferred embodiment, the prostate cancer protein to which antibodies are raised is a transmembrane protein. Without being bound by theory, antibodies used for treatment, bind the extracellular domain of the prostate cancer protein and prevent it from binding to other proteins, such as circulating ligands or cell-associated molecules. The antibody may cause down-regulation of the transmembrane prostate cancer protein. As will be appreciated by one of ordinary skill in the art, the antibody may be a competitive, noncompetitive or uncompetitive inhibitor of protein binding to the extracellular domain of the prostate cancer protein. The antibody is also an antagonist of the prostate cancer protein. Further, the antibody prevents activation of the transmembrane prostate cancer protein. In one aspect, when the antibody prevents the binding of other molecules to the prostate cancer protein, the antibody prevents growth of the cell. The antibody may also be used to target or sensitize the cell to cytotoxic agents, including, but not limited to TNF-α, TNF-β, IL-1, INF-γ and IL-2, or chemotherapeutic agents including 5FU, vinblastine, actinomycin D, cisplatin, methotrexate, and the like. In some instances the antibody belongs to a sub-type that activates serum complement when complexed with the transmembrane protein thereby mediating cytotoxicity or antigen-dependent cytotoxicity (ADCC). Thus, prostate cancer is treated by administering to a patient antibodies directed against the transmembrane prostate cancer protein. Antibody-labeling may activate a co-toxin, localize a toxin payload, or otherwise provide means to locally ablate cells.

In another preferred embodiment, the antibody is conjugated to an effector moiety. The effector moiety can be any number of molecules, including labelling moieties such as radioactive labels or fluorescent labels, or can be a therapeutic moiety. In one aspect the therapeutic moiety is a small molecule that modulates the activity of the prostate cancer protein. In another aspect the therapeutic moiety modulates the activity of molecules associated with or in close proximity to the prostate cancer protein. The therapeutic moiety

may inhibit enzymatic activity such as protease or collagenase or protein kinase activity associated with prostate cancer.

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In a preferred embodiment, the therapeutic moiety can also be a cytotoxic agent. In this method, targeting the cytotoxic agent to prostate cancer tissue or cells, results in a reduction in the number of afflicted cells, thereby reducing symptoms associated with prostate cancer. Cytotoxic agents are numerous and varied and include, but are not limited to, cytotoxic drugs or toxins or active fragments of such toxins. Suitable toxins and their corresponding fragments include diphtheria A chain, exotoxin A chain, ricin A chain, abrin A chain, curcin, crotin, phenomycin, enomycin and the like. Cytotoxic agents also include radiochemicals made by conjugating radioisotopes to antibodies raised against prostate cancer proteins, or binding of a radionuclide to a chelating agent that has been covalently attached to the antibody. Targeting the therapeutic moiety to transmembrane prostate cancer proteins not only serves to increase the local concentration of therapeutic moiety in the prostate cancer afflicted area, but also serves to reduce deleterious side effects that may be associated with the therapeutic moiety.

In another preferred embodiment, the prostate cancer protein against which the antibodies are raised is an intracellular protein. In this case, the antibody may be conjugated to a protein which facilitates entry into the cell. In one case, the antibody enters the cell by endocytosis. In another embodiment, a nucleic acid encoding the antibody is administered to the individual or cell. Moreover, wherein the prostate cancer protein can be targeted within a cell, i.e., the nucleus, an antibody thereto contains a signal for that target localization, i.e., a nuclear localization signal.

The prostate cancer antibodies of the invention specifically bind to prostate cancer proteins. By "specifically bind" herein is meant that the antibodies bind to the protein with a K_d of at least about 0.1 mM, more usually at least about 1 μ M, preferably at least about 0.1 μ M or better, and most preferably, 0.01 μ M or better. Selectivity of binding is also important.

Detection of prostate cancer sequence for diagnostic and therapeutic applications

In one aspect, the RNA expression levels of genes are determined for different cellular states in the prostate cancer phenotype. Expression levels of genes in normal tissue

(i.e., not undergoing prostate cancer) and in prostate cancer tissue (and in some cases, for varying severities of prostate cancer that relate to prognosis, as outlined below) are evaluated to provide expression profiles. An expression profile of a particular cell state or point of development is essentially a "fingerprint" of the state. While two states may have any particular gene similarly expressed, the evaluation of a number of genes simultaneously allows the generation of a gene expression profile that is reflective of the state of the cell. By comparing expression profiles of cells in different states, information regarding which genes are important (including both up- and down-regulation of genes) in each of these states is obtained. Then, diagnosis may be performed or confirmed to determine whether a tissue sample has the gene expression profile of normal or cancerous tissue. This will provide for molecular diagnosis of related conditions.

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"Differential expression," or grammatical equivalents as used herein, refers to qualitative or quantitative differences in the temporal and/or cellular gene expression patterns within and among cells and tissue. Thus, a differentially expressed gene can qualitatively have its expression altered, including an activation or inactivation, in, e.g., normal versus prostate cancer tissue. Genes may be turned on or turned off in a particular state, relative to another state thus permitting comparison of two or more states. A qualitatively regulated gene will exhibit an expression pattern within a state or cell type which is detectable by standard techniques. Some genes will be expressed in one state or cell type, but not in both. Alternatively, the difference in expression may be quantitative, e.g., in that expression is increased or decreased; i.e., gene expression is either upregulated, resulting in an increased amount of transcript, or downregulated, resulting in a decreased amount of transcript. The degree to which expression differs need only be large enough to quantify via standard characterization techniques as outlined below, such as by use of Affymetrix GeneChip[™] expression arrays, Lockhart, *Nature Biotechnology* 14:1675-1680 (1996), hereby expressly incorporated by reference. Other techniques include, but are not limited to, quantitative reverse transcriptase PCR, northern analysis and RNase protection. As outlined above, preferably the change in expression (i.e., upregulation or downregulation) is at least about 50%, more preferably at least about 100%, more preferably at least about 150%, more preferably at least about 200%, with from 300 to at least 1000% being especially preferred.

Evaluation may be at the gene transcript, or the protein level. The amount of gene expression may be monitored using nucleic acid probes to the DNA or RNA equivalent of the gene transcript, and the quantification of gene expression levels, or, alternatively, the final gene product itself (protein) can be monitored, e.g., with antibodies to the prostate cancer protein and standard immunoassays (ELISAs, etc.) or other techniques, including mass spectroscopy assays, 2D gel electrophoresis assays, etc. Proteins corresponding to prostate cancer genes, i.e., those identified as being important in a prostate cancer phenotype, can be evaluated in a prostate cancer diagnostic test.

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In a preferred embodiment, gene expression monitoring is performed simultaneously on a number of genes. Multiple protein expression monitoring can be performed as well. Similarly, these assays may be performed on an individual basis as well.

In this embodiment, the prostate cancer nucleic acid probes are attached to biochips as outlined herein for the detection and quantification of prostate cancer sequences in a particular cell. The assays are further described below in the example. PCR techniques can be used to provide greater sensitivity.

In a preferred embodiment nucleic acids encoding the prostate cancer protein are detected. Although DNA or RNA encoding the prostate cancer protein may be detected, of particular interest are methods wherein an mRNA encoding a prostate cancer protein is detected. Probes to detect mRNA can be a nucleotide/deoxynucleotide probe that is complementary to and hybridizes with the mRNA and includes, but is not limited to, oligonucleotides, cDNA or RNA. Probes also should contain a detectable label, as defined herein. In one method the mRNA is detected after immobilizing the nucleic acid to be examined on a solid support such as nylon membranes and hybridizing the probe with the sample. Following washing to remove the non-specifically bound probe, the label is detected. In another method detection of the mRNA is performed in situ. In this method permeabilized cells or tissue samples are contacted with a detectably labeled nucleic acid probe for sufficient time to allow the probe to hybridize with the target mRNA. Following washing to remove the non-specifically bound probe, the label is detected. For example a digoxygenin labeled riboprobe (RNA probe) that is complementary to the mRNA encoding a prostate cancer protein is detected by binding the digoxygenin with an anti-digoxygenin

secondary antibody and developed with nitro blue tetrazolium and 5-bromo-4-chloro-3-indoyl phosphate.

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In a preferred embodiment, various proteins from the three classes of proteins as described herein (secreted, transmembrane or intracellular proteins) are used in diagnostic assays. The prostate cancer proteins, antibodies, nucleic acids, modified proteins and cells containing prostate cancer sequences are used in diagnostic assays. This can be performed on an individual gene or corresponding polypeptide level. In a preferred embodiment, the expression profiles are used, preferably in conjunction with high throughput screening techniques to allow monitoring for expression profile genes and/or corresponding polypeptides.

As described and defined herein, prostate cancer proteins, including intracellular, transmembrane or secreted proteins, find use as markers of prostate cancer. Detection of these proteins in putative prostate cancer tissue allows for detection or diagnosis of prostate cancer. In one embodiment, antibodies are used to detect prostate cancer proteins. A preferred method separates proteins from a sample by electrophoresis on a gel (typically a denaturing and reducing protein gel, but may be another type of gel, including isoelectric focusing gels and the like). Following separation of proteins, the prostate cancer protein is detected, e.g., by immunoblotting with antibodies raised against the prostate cancer protein. Methods of immunoblotting are well known to those of ordinary skill in the art.

In another preferred method, antibodies to the prostate cancer protein find use in *in situ* imaging techniques, e.g., in histology (e.g., *Methods in Cell Biology: Antibodies in Cell Biology*, volume 37 (Asai, ed. 1993)). In this method cells are contacted with from one to many antibodies to the prostate cancer protein(s). Following washing to remove non-specific antibody binding, the presence of the antibody or antibodies is detected. In one embodiment the antibody is detected by incubating with a secondary antibody that contains a detectable label. In another method the primary antibody to the prostate cancer protein(s) contains a detectable label, e.g. an enzyme marker that can act on a substrate. In another preferred embodiment each one of multiple primary antibodies contains a distinct and detectable label. This method finds particular use in simultaneous screening for a plurality of prostate cancer proteins. As will be appreciated by one of ordinary skill in the art, many other histological imaging techniques are also provided by the invention.

In a preferred embodiment the label is detected in a fluorometer which has the ability to detect and distinguish emissions of different wavelengths. In addition, a fluorescence activated cell sorter (FACS) can be used in the method.

In another preferred embodiment, antibodies find use in diagnosing prostate cancer from blood, serum, plasma, stool, and other samples. Such samples, therefore, are useful as samples to be probed or tested for the presence of prostate cancer proteins.

Antibodies can be used to detect a prostate cancer protein by previously described immunoassay techniques including ELISA, immunoblotting (western blotting), immunoprecipitation, BIACORE technology and the like. Conversely, the presence of antibodies may indicate an immune response against an endogenous prostate cancer protein.

In a preferred embodiment, in situ hybridization of labeled prostate cancer nucleic acid probes to tissue arrays is done. For example, arrays of tissue samples, including prostate cancer tissue and/or normal tissue, are made. In situ hybridization (see, e.g., Ausubel, supra) is then performed. When comparing the fingerprints between an individual and a standard, the skilled artisan can make a diagnosis, a prognosis, or a prediction based on the findings. It is further understood that the genes which indicate the diagnosis may differ from those which indicate the prognosis and molecular profiling of the condition of the cells may lead to distinctions between responsive or refractory conditions or may be predictive of outcomes.

In a preferred embodiment, the prostate cancer proteins, antibodies, nucleic acids, modified proteins and cells containing prostate cancer sequences are used in prognosis assays. As above, gene expression profiles can be generated that correlate to prostate cancer, in terms of long term prognosis. Again, this may be done on either a protein or gene level, with the use of genes being preferred. As above, prostate cancer probes may be attached to biochips for the detection and quantification of prostate cancer sequences in a tissue or patient. The assays proceed as outlined above for diagnosis. PCR method may provide more sensitive and accurate quantification.

Assays for therapeutic compounds

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In a preferred embodiment members of the proteins, nucleic acids, and antibodies as described herein are used in drug screening assays. The prostate cancer

proteins, antibodies, nucleic acids, modified proteins and cells containing prostate cancer sequences are used in drug screening assays or by evaluating the effect of drug candidates on a "gene expression profile" or expression profile of polypeptides. In a preferred embodiment, the expression profiles are used, preferably in conjunction with high throughput screening techniques to allow monitoring for expression profile genes after treatment with a candidate agent (e.g., Zlokarnik, et al., Science 279:84-8 (1998); Heid, Genome Res 6:986-94, 1996).

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In a preferred embodiment, the prostate cancer proteins, antibodies, nucleic acids, modified proteins and cells containing the native or modified prostate cancer proteins are used in screening assays. That is, the present invention provides novel methods for screening for compositions which modulate the prostate cancer phenotype or an identified physiological function of a prostate cancer protein. As above, this can be done on an individual gene level or by evaluating the effect of drug candidates on a "gene expression profile". In a preferred embodiment, the expression profiles are used, preferably in conjunction with high throughput screening techniques to allow monitoring for expression profile genes after treatment with a candidate agent, see Zlokarnik, *supra*.

Having identified the differentially expressed genes herein, a variety of assays may be executed. In a preferred embodiment, assays may be run on an individual gene or protein level. That is, having identified a particular gene as up regulated in prostate cancer, test compounds can be screened for the ability to modulate gene expression or for binding to the prostate cancer protein. "Modulation" thus includes both an increase and a decrease in gene expression. The preferred amount of modulation will depend on the original change of the gene expression in normal versus tissue undergoing prostate cancer, with changes of at least 10%, preferably 50%, more preferably 100-300%, and in some embodiments 300-1000% or greater. Thus, if a gene exhibits a 4-fold increase in prostate cancer tissue compared to normal tissue, a decrease of about four-fold is often desired; similarly, a 10-fold decrease in prostate cancer tissue compared to normal tissue often provides a target value of a 10-fold increase in expression to be induced by the test compound.

The amount of gene expression may be monitored using nucleic acid probes and the quantification of gene expression levels, or, alternatively, the gene product itself can be monitored, e.g., through the use of antibodies to the prostate cancer protein and standard

immunoassays. Proteomics and separation techniques may also allow quantification of expression.

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In a preferred embodiment, gene expression or protein monitoring of a number of entities, i.e., an expression profile, is monitored simultaneously. Such profiles will typically involve a plurality of those entities described herein..

In this embodiment, the prostate cancer nucleic acid probes are attached to biochips as outlined herein for the detection and quantification of prostate cancer sequences in a particular cell. Alternatively, PCR may be used. Thus, a series, e.g., of microtiter plate, may be used with dispensed primers in desired wells. A PCR reaction can then be performed and analyzed for each well.

Expression monitoring can be performed to identify compounds that modify the expression of one or more prostate cancer-associated sequences, e.g., a polynucleotide sequence set out in Tables 1-16. Generally, in a preferred embodiment, a test modulator is added to the cells prior to analysis. Moreover, screens are also provided to identify agents that modulate prostate cancer, modulate prostate cancer proteins, bind to a prostate cancer protein, or interfere with the binding of a prostate cancer protein and an antibody or other binding partner.

The term "test compound" or "drug candidate" or "modulator" or grammatical equivalents as used herein describes any molecule, e.g., protein, oligopeptide, small organic molecule, polysaccharide, polynucleotide, etc., to be tested for the capacity to directly or indirectly alter the prostate cancer phenotype or the expression of a prostate cancer sequence, e.g., a nucleic acid or protein sequence. In preferred embodiments, modulators alter expression profiles, or expression profile nucleic acids or proteins provided herein. In one embodiment, the modulator suppresses a prostate cancer phenotype, e.g. to a normal tissue fingerprint. In another embodiment, a modulator induced a prostate cancer phenotype. Generally, a plurality of assay mixtures are run in parallel with different agent concentrations to obtain a differential response to the various concentrations. Typically, one of these concentrations serves as a negative control, i.e., at zero concentration or below the level of detection.

Drug candidates encompass numerous chemical classes, though typically they are organic molecules, preferably small organic compounds having a molecular weight of

more than 100 and less than about 2,500 daltons. Preferred small molecules are less than 2000, or less than 1500 or less than 1000 or less than 500 D. Candidate agents comprise functional groups necessary for structural interaction with proteins, particularly hydrogen bonding, and typically include at least an amine, carbonyl, hydroxyl or carboxyl group, preferably at least two of the functional chemical groups. The candidate agents often comprise cyclical carbon or heterocyclic structures and/or aromatic or polyaromatic structures substituted with one or more of the above functional groups. Candidate agents are also found among biomolecules including peptides, saccharides, fatty acids, steroids, purines, pyrimidines, derivatives, structural analogs or combinations thereof. Particularly preferred are peptides.

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In one aspect, a modulator will neutralize the effect of a prostate cancer protein. By "neutralize" is meant that activity of a protein is inhibited or blocked and the consequent effect on the cell.

In certain embodiments, combinatorial libraries of potential modulators will be screened for an ability to bind to a prostate cancer polypeptide or to modulate activity. Conventionally, new chemical entities with useful properties are generated by identifying a chemical compound (called a "lead compound") with some desirable property or activity, e.g., inhibiting activity, creating variants of the lead compound, and evaluating the property and activity of those variant compounds. Often, high throughput screening (HTS) methods are employed for such an analysis.

In one preferred embodiment, high throughput screening methods involve providing a library containing a large number of potential therapeutic compounds (candidate compounds). Such "combinatorial chemical libraries" are then screened in one or more assays to identify those library members (particular chemical species or subclasses) that display a desired characteristic activity. The compounds thus identified can serve as conventional "lead compounds" or can themselves be used as potential or actual therapeutics.

A combinatorial chemical library is a collection of diverse chemical compounds generated by either chemical synthesis or biological synthesis by combining a number of chemical "building blocks" such as reagents. For example, a linear combinatorial chemical library, such as a polypeptide (e.g., mutein) library, is formed by combining a set of chemical building blocks called amino acids in every possible way for a given compound

length (i.e., the number of amino acids in a polypeptide compound). Millions of chemical compounds can be synthesized through such combinatorial mixing of chemical building blocks (Gallop *et al.*, *J. Med. Chem.* 37(9):1233-1251 (1994)).

Preparation and screening of combinatorial chemical libraries is well known to those of skill in the art. Such combinatorial chemical libraries include, but are not limited to, 5 peptide libraries (see, e.g., U.S. Patent No. 5,010,175, Furka, Pept. Prot. Res. 37:487-493 (1991), Houghton et al., Nature, 354:84-88 (1991)), peptoids (PCT Publication No WO 91/19735), encoded peptides (PCT Publication WO 93/20242), random bio-oligomers (PCT Publication WO 92/00091), benzodiazepines (U.S. Pat. No. 5,288,514), diversomers such as 10 hydantoins, benzodiazepines and dipeptides (Hobbs et al., Proc. Nat. Acad. Sci. USA 90:6909-6913 (1993)), vinylogous polypeptides (Hagihara et al., J. Amer. Chem. Soc. 114:6568 (1992)), nonpeptidal peptidomimetics with a Beta-D-Glucose scaffolding (Hirschmann et al., J. Amer. Chem. Soc. 114:9217-9218 (1992)), analogous organic syntheses of small compound libraries (Chen et al., J. Amer. Chem. Soc. 116:2661 (1994)), oligocarbamates (Cho, et al., Science 261:1303 (1993)), and/or peptidyl phosphonates 15 (Campbell et al., J. Org. Chem. 59:658 (1994)). See, generally, Gordon et al., J. Med. Chem. 37:1385 (1994), nucleic acid libraries (see, e.g., Strategene, Corp.), peptide nucleic acid libraries (see, e.g., U.S. Patent 5,539,083), antibody libraries (see, e.g., Vaughn et al., Nature Biotechnology 14(3):309-314 (1996), and PCT/US96/10287), carbohydrate libraries (see, e.g., Liang et al., Science 274:1520-1522 (1996), and U.S. Patent No. 5,593,853), and small 20 organic molecule libraries (see, e.g., benzodiazepines, Baum, C&EN, Jan 18, page 33 (1993); isoprenoids, U.S. Patent No. 5,569,588; thiazolidinones and metathiazanones, U.S. Patent No. 5,549,974; pyrrolidines, U.S. Patent Nos. 5,525,735 and 5,519,134; morpholino compounds, U.S. Patent No. 5,506,337; benzodiazepines, U.S. Patent No. 5,288,514; and the like).

Devices for the preparation of combinatorial libraries are commercially available (*see*, *e.g.*, 357 MPS, 390 MPS, Advanced Chem Tech, Louisville KY, Symphony, Rainin, Woburn, MA, 433A Applied Biosystems, Foster City, CA, 9050 Plus, Millipore, Bedford, MA).

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A number of well known robotic systems have also been developed for solution phase chemistries. These systems include automated workstations like the automated synthesis apparatus developed by Takeda Chemical Industries, LTD. (Osaka,

Japan) and many robotic systems utilizing robotic arms (Zymate II, Zymark Corporation, Hopkinton, Mass.; Orca, Hewlett-Packard, Palo Alto, Calif.), which mimic the manual synthetic operations performed by a chemist. Any of the above devices are suitable for use with the present invention. The nature and implementation of modifications to these devices (if any) so that they can operate as discussed herein will be apparent to persons skilled in the relevant art. In addition, numerous combinatorial libraries are themselves commercially available (see, e.g., ComGenex, Princeton, N.J., Asinex, Moscow, Ru, Tripos, Inc., St. Louis, MO, ChemStar, Ltd, Moscow, RU, 3D Pharmaceuticals, Exton, PA, Martek Biosciences, Columbia, MD, etc.).

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The assays to identify modulators are amenable to high throughput screening. Preferred assays thus detect enhancement or inhibition of prostate cancer gene transcription, inhibition or enhancement of polypeptide expression, and inhibition or enhancement of polypeptide activity.

High throughput assays for the presence, absence, quantification, or other properties of particular nucleic acids or protein products are well known to those of skill in the art. Similarly, binding assays and reporter gene assays are similarly well known. Thus, e.g., U.S. Patent No. 5,559,410 discloses high throughput screening methods for proteins, U.S. Patent No. 5,585,639 discloses high throughput screening methods for nucleic acid binding (i.e., in arrays), while U.S. Patent Nos. 5,576,220 and 5,541,061 disclose high throughput methods of screening for ligand/antibody binding.

In addition, high throughput screening systems are commercially available (see, e.g., Zymark Corp., Hopkinton, MA; Air Technical Industries, Mentor, OH; Beckman Instruments, Inc. Fullerton, CA; Precision Systems, Inc., Natick, MA, etc.). These systems typically automate entire procedures, including all sample and reagent pipetting, liquid dispensing, timed incubations, and final readings of the microplate in detector(s) appropriate for the assay. These configurable systems provide high throughput and rapid start up as well as a high degree of flexibility and customization. The manufacturers of such systems provide detailed protocols for various high throughput systems. Thus, e.g., Zymark Corp. provides technical bulletins describing screening systems for detecting the modulation of gene transcription, ligand binding, and the like.

In one embodiment, modulators are proteins, often naturally occurring proteins or fragments of naturally occurring proteins. Thus, e.g., cellular extracts containing proteins, or random or directed digests of proteinaceous cellular extracts, may be used. In this way libraries of proteins may be made for screening in the methods of the invention. Particularly preferred in this embodiment are libraries of bacterial, fungal, viral, and mammalian proteins, with the latter being preferred, and human proteins being especially preferred. Particularly useful test compound will be directed to the class of proteins to which the target belongs, e.g., substrates for enzymes or ligands and receptors.

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In a preferred embodiment, modulators are peptides of from about 5 to about 30 amino acids, with from about 5 to about 20 amino acids being preferred, and from about 7 to about 15 being particularly preferred. The peptides may be digests of naturally occurring proteins as is outlined above, random peptides, or "biased" random peptides. By "randomized" or grammatical equivalents herein is meant that each nucleic acid and peptide consists of essentially random nucleotides and amino acids, respectively. Since generally these random peptides (or nucleic acids, discussed below) are chemically synthesized, they may incorporate any nucleotide or amino acid at any position. The synthetic process can be designed to generate randomized proteins or nucleic acids, to allow the formation of all or most of the possible combinations over the length of the sequence, thus forming a library of randomized candidate bioactive proteinaceous agents.

In one embodiment, the library is fully randomized, with no sequence preferences or constants at any position. In a preferred embodiment, the library is biased. That is, some positions within the sequence are either held constant, or are selected from a limited number of possibilities. For example, in a preferred embodiment, the nucleotides or amino acid residues are randomized within a defined class, e.g., of hydrophobic amino acids, hydrophilic residues, sterically biased (either small or large) residues, towards the creation of nucleic acid binding domains, the creation of cysteines, for cross-linking, prolines for SH-3 domains, serines, threonines, tyrosines or histidines for phosphorylation sites, etc., or to purines, etc.

Modulators of prostate cancer can also be nucleic acids, as defined below. As described above generally for proteins, nucleic acid modulating agents may be naturally occurring nucleic acids, random nucleic acids, or "biased" random nucleic acids. For

example, digests of procaryotic or eucaryotic genomes may be used as is outlined above for proteins.

In certain embodiments, the activity of a prostate cancer-associated protein is down-regulated, or entirely inhibited, by the use of antisense polynucleotide, *i.e.*, a nucleic acid complementary to, and which can preferably hybridize specifically to, a coding mRNA nucleic acid sequence, e.g., a prostate cancer protein mRNA, or a subsequence thereof. Binding of the antisense polynucleotide to the mRNA reduces the translation and/or stability of the mRNA.

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In the context of this invention, antisense polynucleotides can comprise naturally-occurring nucleotides, or synthetic species formed from naturally-occurring subunits or their close homologs. Antisense polynucleotides may also have altered sugar moieties or inter-sugar linkages. Exemplary among these are the phosphorothioate and other sulfur containing species which are known for use in the art. Analogs are comprehended by this invention so long as they function effectively to hybridize with the prostate cancer protein mRNA. See, e.g., Isis Pharmaceuticals, Carlsbad, CA; Sequitor, Inc., Natick, MA.

Such antisense polynucleotides can readily be synthesized using recombinant means, or can be synthesized *in vitro*. Equipment for such synthesis is sold by several vendors, including Applied Biosystems. The preparation of other oligonucleotides such as phosphorothioates and alkylated derivatives is also well known to those of skill in the art.

Antisense molecules as used herein include antisense or sense oligonucleotides. Sense oligonucleotides can, e.g., be employed to block transcription by binding to the anti-sense strand. The antisense and sense oligonucleotide comprise a single-stranded nucleic acid sequence (either RNA or DNA) capable of binding to target mRNA (sense) or DNA (antisense) sequences for prostate cancer molecules. Antisense or sense oligonucleotides, according to the present invention, comprise a fragment generally at least about 14 nucleotides, preferably from about 14 to 30 nucleotides. The ability to derive an antisense or a sense oligonucleotide, based upon a cDNA sequence encoding a given protein is described in, e.g., Stein & Cohen (Cancer Res. 48:2659 (1988 and van der Krol et al. (BioTechniques 6:958 (1988)).

In addition to antisense polynucleotides, ribozymes can be used to target and inhibit transcription of prostate cancer-associated nucleotide sequences. A ribozyme is an

RNA molecule that catalytically cleaves other RNA molecules. Different kinds of ribozymes have been described, including group I ribozymes, hammerhead ribozymes, hairpin ribozymes, RNase P, and axhead ribozymes (see, e.g., Castanotto et al., Adv. in Pharmacology 25: 289-317 (1994) for a general review of the properties of different ribozymes).

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The general features of hairpin ribozymes are described, e.g., in Hampel et al., Nucl. Acids Res. 18:299-304 (1990); European Patent Publication No. 0 360 257; U.S. Patent No. 5,254,678. Methods of preparing are well known to those of skill in the art (see, e.g., WO 94/26877; Ojwang et al., Proc. Natl. Acad. Sci. USA 90:6340-6344 (1993); Yamada et al., Human Gene Therapy 1:39-45 (1994); Leavitt et al., Proc. Natl. Acad. Sci. USA 92:699-703 (1995); Leavitt et al., Human Gene Therapy 5:1151-120 (1994); and Yamada et al., Virology 205: 121-126 (1994)).

Polynucleotide modulators of prostate cancer may be introduced into a cell containing the target nucleotide sequence by formation of a conjugate with a ligand binding molecule, as described in WO 91/04753. Suitable ligand binding molecules include, but are not limited to, cell surface receptors, growth factors, other cytokines, or other ligands that bind to cell surface receptors. Preferably, conjugation of the ligand binding molecule does not substantially interfere with the ability of the ligand binding molecule to bind to its corresponding molecule or receptor, or block entry of the sense or antisense oligonucleotide or its conjugated version into the cell. Alternatively, a polynucleotide modulator of prostate cancer may be introduced into a cell containing the target nucleic acid sequence, e.g., by formation of an polynucleotide-lipid complex, as described in WO 90/10448. It is understood that the use of antisense molecules or knock out and knock in models may also be used in screening assays as discussed above, in addition to methods of treatment.

As noted above, gene expression monitoring is conveniently used to test candidate modultors (e.g., protein, nucleic acid or small molecule). After the candidate agent has been added and the cells allowed to incubate for some period of time, the sample containing a target sequence to be analyzed is added to the biochip. If required, the target sequence is prepared using known techniques. For example, the sample may be treated to lyse the cells, using known lysis buffers, electroporation, etc., with purification and/or amplification such as PCR performed as appropriate. For example, an in vitro transcription

with labels covalently attached to the nucleotides is performed. Generally, the nucleic acids are labeled with biotin-FITC or PE, or with cy3 or cy5.

In a preferred embodiment, the target sequence is labeled with, e.g., a fluorescent, a chemiluminescent, a chemical, or a radioactive signal, to provide a means of detecting the target sequence's specific binding to a probe. The label also can be an enzyme, such as, alkaline phosphatase or horseradish peroxidase, which when provided with an appropriate substrate produces a product that can be detected. Alternatively, the label can be a labeled compound or small molecule, such as an enzyme inhibitor, that binds but is not catalyzed or altered by the enzyme. The label also can be a moiety or compound, such as, an epitope tag or biotin which specifically binds to streptavidin. For the example of biotin, the streptavidin is labeled as described above, thereby, providing a detectable signal for the bound target sequence. Unbound labeled streptavidin is typically removed prior to analysis.

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As will be appreciated by those in the art, these assays can be direct hybridization assays or can comprise "sandwich assays", which include the use of multiple probes, as is generally outlined in U.S. Patent Nos. 5,681,702, 5,597,909, 5,545,730, 5,594,117, 5,591,584, 5,571,670, 5,580,731, 5,571,670, 5,591,584, 5,624,802, 5,635,352, 5,594,118, 5,359,100, 5,124,246 and 5,681,697, all of which are hereby incorporated by reference. In this embodiment, in general, the target nucleic acid is prepared as outlined above, and then added to the biochip comprising a plurality of nucleic acid probes, under conditions that allow the formation of a hybridization complex.

A variety of hybridization conditions may be used in the present invention, including high, moderate and low stringency conditions as outlined above. The assays are generally run under stringency conditions which allows formation of the label probe hybridization complex only in the presence of target. Stringency can be controlled by altering a step parameter that is a thermodynamic variable, including, but not limited to, temperature, formamide concentration, salt concentration, chaotropic salt concentration pH, organic solvent concentration, etc.

These parameters may also be used to control non-specific binding, as is generally outlined in U.S. Patent No. 5,681,697. Thus it may be desirable to perform certain steps at higher stringency conditions to reduce non-specific binding.

The reactions outlined herein may be accomplished in a variety of ways. Components of the reaction may be added simultaneously, or sequentially, in different orders, with preferred embodiments outlined below. In addition, the reaction may include a variety of other reagents. These include salts, buffers, neutral proteins, e.g. albumin, detergents, etc. which may be used to facilitate optimal hybridization and detection, and/or reduce non-specific or background interactions. Reagents that otherwise improve the efficiency of the assay, such as protease inhibitors, nuclease inhibitors, anti-microbial agents, etc., may also be used as appropriate, depending on the sample preparation methods and purity of the target.

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The assay data are analyzed to determine the expression levels, and changes in expression levels as between states, of individual genes, forming a gene expression profile.

Screens are performed to identify modulators of the prostate cancer phenotype. In one embodiment, screening is performed to identify modulators that can induce or suppress a particular expression profile, thus preferably generating the associated phenotype. In another embodiment, e.g., for diagnostic applications, having identified differentially expressed genes important in a particular state, screens can be performed to identify modulators that alter expression of individual genes. In an another embodiment, screening is performed to identify modulators that alter a biological function of the expression product of a differentially expressed gene. Again, having identified the importance of a gene in a particular state, screens are performed to identify agents that bind and/or modulate the biological activity of the gene product.

In addition screens can be done for genes that are induced in response to a candidate agent. After identifying a modulator based upon its ability to suppress a prostate cancer expression pattern leading to a normal expression pattern, or to modulate a single prostate cancer gene expression profile so as to mimic the expression of the gene from normal tissue, a screen as described above can be performed to identify genes that are specifically modulated in response to the agent. Comparing expression profiles between normal tissue and agent treated prostate cancer tissue reveals genes that are not expressed in normal tissue or prostate cancer tissue, but are expressed in agent treated tissue. These agent-specific sequences can be identified and used by methods described herein for prostate cancer genes or proteins. In particular these sequences and the proteins they encode find use in marking or identifying agent treated cells. In addition, antibodies can be raised against the

agent induced proteins and used to target novel therapeutics to the treated prostate cancer tissue sample.

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Thus, in one embodiment, a test compound is administered to a population of prostate cancer cells, that have an associated prostate cancer expression profile. By "administration" or "contacting" herein is meant that the candidate agent is added to the cells in such a manner as to allow the agent to act upon the cell, whether by uptake and intracellular action, or by action at the cell surface. In some embodiments, nucleic acid encoding a proteinaceous candidate agent (i.e., a peptide) may be put into a viral construct such as an adenoviral or retroviral construct, and added to the cell, such that expression of the peptide agent is accomplished, e.g., PCT US97/01019. Regulatable gene therapy systems can also be used.

Once the test compound has been administered to the cells, the cells can be washed if desired and are allowed to incubate under preferably physiological conditions for some period of time. The cells are then harvested and a new gene expression profile is generated, as outlined herein.

Thus, e.g., prostate cancer tissue may be screened for agents that modulate, e.g., induce or suppress the prostate cancer phenotype. A change in at least one gene, preferably many, of the expression profile indicates that the agent has an effect on prostate cancer activity. By defining such a signature for the prostate cancer phenotype, screens for new drugs that alter the phenotype can be devised. With this approach, the drug target need not be known and need not be represented in the original expression screening platform, nor does the level of transcript for the target protein need to change.

In a preferred embodiment, as outlined above, screens may be done on individual genes and gene products (proteins). That is, having identified a particular differentially expressed gene as important in a particular state, screening of modulators of either the expression of the gene or the gene product itself can be done. The gene products of differentially expressed genes are sometimes referred to herein as "prostate cancer proteins" or a "prostate cancer modulatory protein". The prostate cancer modulatory protein may be a fragment, or alternatively, be the full length protein to the fragment encoded by the nucleic acids of Tables 1-16. Preferably, the prostate cancer modulatory protein is a fragment. In a preferred embodiment, the prostate cancer amino acid sequence which is used to determine

sequence identity or similarity is encoded by a nucleic acid of Tables 1-16. In another embodiment, the sequences are naturally occurring allelic variants of a protein encoded by a nucleic acid of Tables 1-16. In another embodiment, the sequences are sequence variants as further described herein.

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Preferably, the prostate cancer modulatory protein is a fragment of approximately 14 to 24 amino acids long. More preferably the fragment is a soluble fragment. Preferably, the fragment includes a non-transmembrane region. In a preferred embodiment, the fragment has an N-terminal Cys to aid in solubility. In one embodiment, the C-terminus of the fragment is kept as a free acid and the N-terminus is a free amine to aid in coupling, i.e., to cysteine.

In one embodiment the prostate cancer proteins are conjugated to an immunogenic agent as discussed herein. In one embodiment the prostate cancer protein is conjugated to BSA.

Measurements of prostate cancer polypeptide activity, or of prostate cancer or the prostate cancer phenotype can be performed using a variety of assays. For example, the effects of the test compounds upon the function of the prostate cancer polypeptides can be measured by examining parameters described above. A suitable physiological change that affects activity can be used to assess the influence of a test compound on the polypeptides of this invention. When the functional consequences are determined using intact cells or animals, one can also measure a variety of effects such as, in the case of prostate cancer associated with tumors, tumor growth, tumor metastasis, neovascularization, hormone release, transcriptional changes to both known and uncharacterized genetic markers (e.g., northern blots), changes in cell metabolism such as cell growth or pH changes, and changes in intracellular second messengers such as cGMP. In the assays of the invention, mammalian prostate cancer polypeptide is typically used, e.g., mouse, preferably human.

Assays to identify compounds with modulating activity can be performed in vitro. For example, a prostate cancer polypeptide is first contacted with a potential modulator and incubated for a suitable amount of time, e.g., from 0.5 to 48 hours. In one embodiment, the prostate cancer polypeptide levels are determined in vitro by measuring the level of protein or mRNA. The level of protein is measured using immunoassays such as western blotting, ELISA and the like with an antibody that selectively binds to the prostate cancer

polypeptide or a fragment thereof. For measurement of mRNA, amplification, e.g., using PCR, LCR, or hybridization assays, e.g., northern hybridization, RNAse protection, dot blotting, are preferred. The level of protein or mRNA is detected using directly or indirectly labeled detection agents, e.g., fluorescently or radioactively labeled nucleic acids, radioactively or enzymatically labeled antibodies, and the like, as described herein.

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Alternatively, a reporter gene system can be devised using the prostate cancer protein promoter operably linked to a reporter gene such as luciferase, green fluorescent protein, CAT, or β -gal. The reporter construct is typically transfected into a cell. After treatment with a potential modulator, the amount of reporter gene transcription, translation, or activity is measured according to standard techniques known to those of skill in the art.

In a preferred embodiment, as outlined above, screens may be done on individual genes and gene products (proteins). That is, having identified a particular differentially expressed gene as important in a particular state, screening of modulators of the expression of the gene or the gene product itself can be done. The gene products of differentially expressed genes are sometimes referred to herein as "prostate cancer proteins." The prostate cancer protein may be a fragment, or alternatively, be the full length protein to a fragment shown herein.

In one embodiment, screening for modulators of expression of specific genes is performed. Typically, the expression of only one or a few genes are evaluated. In another embodiment, screens are designed to first find compounds that bind to differentially expressed proteins. These compounds are then evaluated for the ability to modulate differentially expressed activity. Moreover, once initial candidate compounds are identified, variants can be further screened to better evaluate structure activity relationships.

In a preferred embodiment, binding assays are done. In general, purified or isolated gene product is used; that is, the gene products of one or more differentially expressed nucleic acids are made. For example, antibodies are generated to the protein gene products, and standard immunoassays are run to determine the amount of protein present. Alternatively, cells comprising the prostate cancer proteins can be used in the assays.

Thus, in a preferred embodiment, the methods comprise combining a prostate cancer protein and a candidate compound, and determining the binding of the compound to the prostate cancer protein. Preferred embodiments utilize the human prostate cancer protein,

although other mammalian proteins may also be used, e.g. for the development of animal models of human disease. In some embodiments, as outlined herein, variant or derivative prostate cancer proteins may be used.

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Generally, in a preferred embodiment of the methods herein, the prostate cancer protein or the candidate agent is non-diffusably bound to an insoluble support having isolated sample receiving areas (e.g. a microtiter plate, an array, etc.). The insoluble supports may be made of any composition to which the compositions can be bound, is readily separated from soluble material, and is otherwise compatible with the overall method of screening. The surface of such supports may be solid or porous and of any convenient shape. Examples of suitable insoluble supports include microtiter plates, arrays, membranes and beads. These are typically made of glass, plastic (e.g., polystyrene), polysaccharides, nylon or nitrocellulose, teflonTM, etc. Microtiter plates and arrays are especially convenient because a large number of assays can be carried out simultaneously, using small amounts of reagents and samples. The particular manner of binding of the composition is not crucial so long as it is compatible with the reagents and overall methods of the invention, maintains the activity of the composition and is nondiffusable. Preferred methods of binding include the use of antibodies (which do not sterically block either the ligand binding site or activation sequence when the protein is bound to the support), direct binding to "sticky" or ionic supports, chemical crosslinking, the synthesis of the protein or agent on the surface, etc. Following binding of the protein or agent, excess unbound material is removed by washing. The sample receiving areas may then be blocked through incubation with bovine serum albumin (BSA), casein or other innocuous protein or other moiety.

In a preferred embodiment, the prostate cancer protein is bound to the support, and a test compound is added to the assay. Alternatively, the candidate agent is bound to the support and the prostate cancer protein is added. Novel binding agents include specific antibodies, non-natural binding agents identified in screens of chemical libraries, peptide analogs, etc. Of particular interest are screening assays for agents that have a low toxicity for human cells. A wide variety of assays may be used for this purpose, including labeled in vitro protein-protein binding assays, electrophoretic mobility shift assays, immunoassays for protein binding, functional assays (phosphorylation assays, etc.) and the like.

The determination of the binding of the test modulating compound to the prostate cancer protein may be done in a number of ways. In a preferred embodiment, the compound is labeled, and binding determined directly, e.g., by attaching all or a portion of the prostate cancer protein to a solid support, adding a labeled candidate agent (e.g., a fluorescent label), washing off excess reagent, and determining whether the label is present on the solid support. Various blocking and washing steps may be utilized as appropriate.

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In some embodiments, only one of the components is labeled, e.g., the proteins (or proteinaceous candidate compounds) can be labeled. Alternatively, more than one component can be labeled with different labels, e.g., ¹²⁵I for the proteins and a fluorophor for the compound. Proximity reagents, e.g., quenching or energy transfer reagents are also useful.

In one embodiment, the binding of the test compound is determined by competitive binding assay. The competitor is a binding moiety known to bind to the target molecule (i.e., a prostate cancer protein), such as an antibody, peptide, binding partner, ligand, etc. Under certain circumstances, there may be competitive binding between the compound and the binding moiety, with the binding moiety displacing the compound. In one embodiment, the test compound is labeled. Either the compound, or the competitor, or both, is added first to the protein for a time sufficient to allow binding, if present. Incubations may be performed at a temperature which facilitates optimal activity, typically between 4 and 40°C. Incubation periods are typically optimized, e.g., to facilitate rapid high throughput screening. Typically between 0.1 and 1 hour will be sufficient. Excess reagent is generally removed or washed away. The second component is then added, and the presence or absence of the labeled component is followed, to indicate binding.

In a preferred embodiment, the competitor is added first, followed by the test compound. Displacement of the competitor is an indication that the test compound is binding to the prostate cancer protein and thus is capable of binding to, and potentially modulating, the activity of the prostate cancer protein. In this embodiment, either component can be labeled. Thus, e.g., if the competitor is labeled, the presence of label in the wash solution indicates displacement by the agent. Alternatively, if the test compound is labeled, the presence of the label on the support indicates displacement.

In an alternative embodiment, the test compound is added first, with incubation and washing, followed by the competitor. The absence of binding by the competitor may indicate that the test compound is bound to the prostate cancer protein with a higher affinity. Thus, if the test compound is labeled, the presence of the label on the support, coupled with a lack of competitor binding, may indicate that the test compound is capable of binding to the prostate cancer protein.

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In a preferred embodiment, the methods comprise differential screening to identity agents that are capable of modulating the activity of the prostate cancer proteins. In this embodiment, the methods comprise combining a prostate cancer protein and a competitor in a first sample. A second sample comprises a test compound, a prostate cancer protein, and a competitor. The binding of the competitor is determined for both samples, and a change, or difference in binding between the two samples indicates the presence of an agent capable of binding to the prostate cancer protein and potentially modulating its activity. That is, if the binding of the competitor is different in the second sample relative to the first sample, the agent is capable of binding to the prostate cancer protein.

Alternatively, differential screening is used to identify drug candidates that bind to the native prostate cancer protein, but cannot bind to modified prostate cancer proteins. The structure of the prostate cancer protein may be modeled, and used in rational drug design to synthesize agents that interact with that site. Drug candidates that affect the activity of a prostate cancer protein are also identified by screening drugs for the ability to either enhance or reduce the activity of the protein.

Positive controls and negative controls may be used in the assays. Preferably control and test samples are performed in at least triplicate to obtain statistically significant results. Incubation of all samples is for a time sufficient for the binding of the agent to the protein. Following incubation, samples are washed free of non-specifically bound material and the amount of bound, generally labeled agent determined. For example, where a radiolabel is employed, the samples may be counted in a scintillation counter to determine the amount of bound compound.

A variety of other reagents may be included in the screening assays. These include reagents like salts, neutral proteins, e.g. albumin, detergents, etc. which may be used to facilitate optimal protein-protein binding and/or reduce non-specific or background

interactions. Also reagents that otherwise improve the efficiency of the assay, such as protease inhibitors, nuclease inhibitors, anti-microbial agents, etc., may be used. The mixture of components may be added in an order that provides for the requisite binding.

In a preferred embodiment, the invention provides methods for screening for a compound capable of modulating the activity of a prostate cancer protein. The methods comprise adding a test compound, as defined above, to a cell comprising prostate cancer proteins. Preferred cell types include almost any cell. The cells contain a recombinant nucleic acid that encodes a prostate cancer protein. In a preferred embodiment, a library of candidate agents are tested on a plurality of cells.

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In one aspect, the assays are evaluated in the presence or absence or previous or subsequent exposure of physiological signals, e.g. hormones, antibodies, peptides, antigens, cytokines, growth factors, action potentials, pharmacological agents including chemotherapeutics, radiation, carcinogenics, or other cells (i.e. cell-cell contacts). In another example, the determinations are determined at different stages of the cell cycle process.

In this way, compounds that modulate prostate cancer agents are identified. Compounds with pharmacological activity are able to enhance or interfere with the activity of the prostate cancer protein. Once identified, similar structures are evaluated to identify critical structural feature of the compound.

In one embodiment, a method of inhibiting prostate cancer cell division is provided. The method comprises administration of a prostate cancer inhibitor. In another embodiment, a method of inhibiting prostate cancer is provided. The method comprises administration of a prostate cancer inhibitor. In a further embodiment, methods of treating cells or individuals with prostate cancer are provided. The method comprises administration of a prostate cancer inhibitor.

In one embodiment, a prostate cancer inhibitor is an antibody as discussed above. In another embodiment, the prostate cancer inhibitor is an antisense molecule.

A variety of cell growth, proliferation, and metastasis assays are known to those of skill in the art, as described below.

Soft agar growth or colony formation in suspension

Normal cells require a solid substrate to attach and grow. When the cells are transformed, they lose this phenotype and grow detached from the substrate. For example,

transformed cells can grow in stirred suspension culture or suspended in semi-solid media, such as semi-solid or soft agar. The transformed cells, when transfected with tumor suppressor genes, regenerate normal phenotype and require a solid substrate to attach and grow. Soft agar growth or colony formation in suspension assays can be used to identify modulators of prostate cancer sequences, which when expressed in host cells, inhibit abnormal cellular proliferation and transformation. A therapeutic compound would reduce or eliminate the host cells' ability to grow in stirred suspension culture or suspended in semi-solid media, such as semi-solid or soft.

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Techniques for soft agar growth or colony formation in suspension assays are described in Freshney, *Culture of Animal Cells a Manual of Basic Technique* (3rd ed., 1994), herein incorporated by reference. *See also*, the methods section of Garkavtsev *et al.* (1996), *supra*, herein incorporated by reference.

Contact inhibition and density limitation of growth

Normal cells typically grow in a flat and organized pattern in a petri dish until they touch other cells. When the cells touch one another, they are contact inhibited and stop growing. When cells are transformed, however, the cells are not contact inhibited and continue to grow to high densities in disorganized foci. Thus, the transformed cells grow to a higher saturation density than normal cells. This can be detected morphologically by the formation of a disoriented monolayer of cells or rounded cells in foci within the regular pattern of normal surrounding cells. Alternatively, labeling index with (³H)-thymidine at saturation density can be used to measure density limitation of growth. See Freshney (1994), supra. The transformed cells, when transfected with tumor suppressor genes, regenerate a normal phenotype and become contact inhibited and would grow to a lower density.

In this assay, labeling index with (³H)-thymidine at saturation density is a preferred method of measuring density limitation of growth. Transformed host cells are transfected with a prostate cancer-associated sequence and are grown for 24 hours at saturation density in non-limiting medium conditions. The percentage of cells labeling with (³H)-thymidine is determined autoradiographically. *See*, Freshney (1994), *supra*.

Growth factor or serum dependence

Transformed cells have a lower serum dependence than their normal counterparts (see, e.g., Temin, J. Natl. Cancer Insti. 37:167-175 (1966); Eagle et al., J. Exp. Med. 131:836-879 (1970)); Freshney, supra. This is in part due to release of various growth factors by the transformed cells. Growth factor or serum dependence of transformed host cells can be compared with that of control.

Tumor specific markers levels

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Tumor cells release an increased amount of certain factors (hereinafter "tumor specific markers") than their normal counterparts. For example, plasminogen activator (PA) is released from human glioma at a higher level than from normal brain cells (see, e.g., Gullino, Angiogenesis, tumor vascularization, and potential interference with tumor growth. in Biological Responses in Cancer, pp. 178-184 (Mihich (ed.) 1985)). Similarly, Tumor angiogenesis factor (TAF) is released at a higher level in tumor cells than their normal counterparts. See, e.g., Folkman, Angiogenesis and Cancer, Sem Cancer Biol. (1992)).

Various techniques which measure the release of these factors are described in Freshney (1994), supra. Also, see, Unkless et al., J. Biol. Chem. 249:4295-4305 (1974); Strickland & Beers, J. Biol. Chem. 251:5694-5702 (1976); Whur et al., Br. J. Cancer 42:305-312 (1980); Gullino, Angiogenesis, tumor vascularization, and potential interference with tumor growth. in Biological Responses in Cancer, pp. 178-184 (Mihich (ed.) 1985); Freshney Anticancer Res. 5:111-130 (1985).

Invasiveness into Matrigel

The degree of invasiveness into Matrigel or some other extracellular matrix constituent can be used as an assay to identify compounds that modulate prostate cancer-associated sequences. Tumor cells exhibit a good correlation between malignancy and invasiveness of cells into Matrigel or some other extracellular matrix constituent. In this assay, tumorigenic cells are typically used as host cells. Expression of a tumor suppressor gene in these host cells would decrease invasiveness of the host cells.

Techniques described in Freshney (1994), *supra*, can be used. Briefly, the level of invasion of host cells can be measured by using filters coated with Matrigel or some

other extracellular matrix constituent. Penetration into the gel, or through to the distal side of the filter, is rated as invasiveness, and rated histologically by number of cells and distance moved, or by prelabeling the cells with ¹²⁵I and counting the radioactivity on the distal side of the filter or bottom of the dish. See, e.g., Freshney (1984), supra.

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Tumor growth in vivo

Effects of prostate cancer-associated sequences on cell growth can be tested in transgenic or immune-suppressed mice. Knock-out transgenic mice can be made, in which the prostate cancer gene is disrupted or in which a prostate cancer gene is inserted. Knock-out transgenic mice can be made by insertion of a marker gene or other heterologous gene into the endogenous prostate cancer gene site in the mouse genome via homologous recombination. Such mice can also be made by substituting the endogenous prostate cancer gene with a mutated version of the prostate cancer gene, or by mutating the endogenous prostate cancer gene, e.g., by exposure to carcinogens.

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A DNA construct is introduced into the nuclei of embryonic stem cells. Cells containing the newly engineered genetic lesion are injected into a host mouse embryo, which is re-implanted into a recipient female. Some of these embryos develop into chimeric mice that possess germ cells partially derived from the mutant cell line. Therefore, by breeding the chimeric mice it is possible to obtain a new line of mice containing the introduced genetic lesion (see, e.g., Capecchi et al., Science 244:1288 (1989)). Chimeric targeted mice can be derived according to Hogan et al., Manipulating the Mouse Embryo: A Laboratory Manual, Cold Spring Harbor Laboratory (1988) and Teratocarcinomas and Embryonic Stem Cells: A Practical Approach, Robertson, ed., IRL Press, Washington, D.C., (1987).

Alternatively, various immune-suppressed or immune-deficient host animals can be used. For example, genetically athymic "nude" mouse (see, e.g., Giovanella et al., J. Natl. Cancer Inst. 52:921 (1974)), a SCID mouse, a thymectomized mouse, or an irradiated mouse (see, e.g., Bradley et al., Br. J. Cancer 38:263 (1978); Selby et al., Br. J. Cancer 41:52 (1980)) can be used as a host. Transplantable tumor cells (typically about 10⁶ cells) injected into isogenic hosts will produce invasive tumors in a high proportions of cases, while normal cells of similar origin will not. In hosts which developed invasive tumors, cells expressing a prostate cancer-associated sequences are injected subcutaneously. After a

suitable length of time, preferably 4-8 weeks, tumor growth is measured (e.g., by volume or by its two largest dimensions) and compared to the control. Tumors that have statistically significant reduction (using, e.g., Student's T test) are said to have inhibited growth.

5 Methods of identifying variant prostate cancer-associated sequences

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Without being bound by theory, expression of various prostate cancer sequences is correlated with prostate cancer. Accordingly, disorders based on mutant or variant prostate cancer genes may be determined. In one embodiment, the invention provides methods for identifying cells containing variant prostate cancer genes, e.g., determining all or part of the sequence of at least one endogenous prostate cancer genes in a cell. This may be accomplished using any number of sequencing techniques. In a preferred embodiment, the invention provides methods of identifying the prostate cancer genetype of an individual, e.g., determining all or part of the sequence of at least one prostate cancer gene of the individual. This is generally done in at least one tissue of the individual, and may include the evaluation of a number of tissues or different samples of the same tissue. The method may include comparing the sequence of the sequenced prostate cancer gene to a known prostate cancer gene, i.e., a wild-type gene.

The sequence of all or part of the prostate cancer gene can then be compared to the sequence of a known prostate cancer gene to determine if any differences exist. This can be done using any number of known homology programs, such as Bestfit, etc. In a preferred embodiment, the presence of a difference in the sequence between the prostate cancer gene of the patient and the known prostate cancer gene correlates with a disease state or a propensity for a disease state, as outlined herein.

In a preferred embodiment, the prostate cancer genes are used as probes to determine the number of copies of the prostate cancer gene in the genome.

In another preferred embodiment, the prostate cancer genes are used as probes to determine the chromosomal localization of the prostate cancer genes. Information such as chromosomal localization finds use in providing a diagnosis or prognosis in particular when chromosomal abnormalities such as translocations, and the like are identified in the prostate cancer gene locus.

Administration of pharmaceutical and vaccine compositions

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In one embodiment, a therapeutically effective dose of a prostate cancer protein or modulator thereof, is administered to a patient. By "therapeutically effective dose" herein is meant a dose that produces effects for which it is administered. The exact dose will depend on the purpose of the treatment, and will be ascertainable by one skilled in the art using known techniques (e.g., Ansel et al., Pharmaceutical Dosage Forms and Drug Delivery; Lieberman, Pharmaceutical Dosage Forms (vols. 1-3, 1992), Dekker, ISBN 0824770846, 082476918X, 0824712692, 0824716981; Lloyd, The Art, Science and Technology of Pharmaceutical Compounding (1999); and Pickar, Dosage Calculations (1999)). As is known in the art, adjustments for prostate cancer degradation, systemic versus localized delivery, and rate of new protease synthesis, as well as the age, body weight, general health, sex, diet, time of administration, drug interaction and the severity of the condition may be necessary, and will be ascertainable with routine experimentation by those skilled in the art. U.S. Patent Application N. 09/687,576, further discloses the use of compositions and methods of diagnosis and treatment in prostate cancer is hereby expressly incorporated by reference.

A "patient" for the purposes of the present invention includes both humans and other animals, particularly mammals. Thus the methods are applicable to both human therapy and veterinary applications. In the preferred embodiment the patient is a mammal, preferably a primate, and in the most preferred embodiment the patient is human.

The administration of the prostate cancer proteins and modulators thereof of the present invention can be done in a variety of ways as discussed above, including, but not limited to, orally, subcutaneously, intravenously, intranasally, transdermally, intraperitoneally, intramuscularly, intrapulmonary, vaginally, rectally, or intraocularly. In some instances, e.g., in the treatment of wounds and inflammation, the prostate cancer proteins and modulators may be directly applied as a solution or spray.

The pharmaceutical compositions of the present invention comprise a prostate cancer protein in a form suitable for administration to a patient. In the preferred embodiment, the pharmaceutical compositions are in a water soluble form, such as being present as pharmaceutically acceptable salts, which is meant to include both acid and base addition salts. "Pharmaceutically acceptable acid addition salt" refers to those salts that retain the

biological effectiveness of the free bases and that are not biologically or otherwise undesirable, formed with inorganic acids such as hydrochloric acid, hydrobromic acid, sulfuric acid, nitric acid, phosphoric acid and the like, and organic acids such as acetic acid, propionic acid, glycolic acid, pyruvic acid, oxalic acid, maleic acid, malonic acid, succinic acid, fumaric acid, tartaric acid, citric acid, benzoic acid, cinnamic acid, mandelic acid, methanesulfonic acid, ethanesulfonic acid, p-toluenesulfonic acid, salicylic acid and the like. "Pharmaceutically acceptable base addition salts" include those derived from inorganic bases such as sodium, potassium, lithium, ammonium, calcium, magnesium, iron, zinc, copper, manganese, aluminum salts and the like. Particularly preferred are the ammonium, potassium, sodium, calcium, and magnesium salts. Salts derived from pharmaceutically acceptable organic non-toxic bases include salts of primary, secondary, and tertiary amines, substituted amines including naturally occurring substituted amines, cyclic amines and basic ion exchange resins, such as isopropylamine, trimethylamine, diethylamine, triethylamine, tripropylamine, and ethanolamine.

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The pharmaceutical compositions may also include one or more of the following: carrier proteins such as serum albumin; buffers; fillers such as microcrystalline cellulose, lactose, corn and other starches; binding agents; sweeteners and other flavoring agents; coloring agents; and polyethylene glycol.

The pharmaceutical compositions can be administered in a variety of unit dosage forms depending upon the method of administration. For example, unit dosage forms suitable for oral administration include, but are not limited to, powder, tablets, pills, capsules and lozenges. It is recognized that prostate cancer protein modulators (e.g., antibodies, antisense constructs, ribozymes, small organic molecules, etc.) when administered orally, should be protected from digestion. This is typically accomplished either by complexing the molecule(s) with a composition to render it resistant to acidic and enzymatic hydrolysis, or by packaging the molecule(s) in an appropriately resistant carrier, such as a liposome or a protection barrier. Means of protecting agents from digestion are well known in the art.

The compositions for administration will commonly comprise a prostate cancer protein modulator dissolved in a pharmaceutically acceptable carrier, preferably an aqueous carrier. A variety of aqueous carriers can be used, e.g., buffered saline and the like. These solutions are sterile and generally free of undesirable matter. These compositions may

be sterilized by conventional, well known sterilization techniques. The compositions may contain pharmaceutically acceptable auxiliary substances as required to approximate physiological conditions such as pH adjusting and buffering agents, toxicity adjusting agents and the like, e.g., sodium acetate, sodium chloride, potassium chloride, calcium chloride, sodium lactate and the like. The concentration of active agent in these formulations can vary widely, and will be selected primarily based on fluid volumes, viscosities, body weight and the like in accordance with the particular mode of administration selected and the patient's needs (e.g., Remington's Pharmaceutical Science (15th ed., 1980) and Goodman & Gillman, The Pharmacologial Basis of Therapeutics (Hardman et al., eds., 1996)).

Thus, a typical pharmaceutical composition for intravenous administration would be about 0.1 to 10 mg per patient per day. Dosages from 0.1 up to about 100 mg per patient per day may be used, particularly when the drug is administered to a secluded site and not into the blood stream, such as into a body cavity or into a lumen of an organ. Substantially higher dosages are possible in topical administration. Actual methods for preparing parenterally administrable compositions will be known or apparent to those skilled in the art, e.g., Remington's Pharmaceutical Science and Goodman and Gillman, The Pharmacologial Basis of Therapeutics, supra.

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The compositions containing modulators of prostate cancer proteins can be administered for therapeutic or prophylactic treatments. In therapeutic applications, compositions are administered to a patient suffering from a disease (e.g., a cancer) in an amount sufficient to cure or at least partially arrest the disease and its complications. An amount adequate to accomplish this is defined as a "therapeutically effective dose." Amounts effective for this use will depend upon the severity of the disease and the general state of the patient's health. Single or multiple administrations of the compositions may be administered depending on the dosage and frequency as required and tolerated by the patient. In any event, the composition should provide a sufficient quantity of the agents of this invention to effectively treat the patient. An amount of modulator that is capable of preventing or slowing the development of cancer in a mammal is referred to as a "prophylactically effective dose." The particular dose required for a prophylactic treatment will depend upon the medical condition and history of the mammal, the particular cancer being prevented, as well as other factors such as age, weight, gender, administration route, efficiency, etc. Such prophylactic

treatments may be used, e.g., in a mammal who has previously had cancer to prevent a recurrence of the cancer, or in a mammal who is suspected of having a significant likelihood of developing cancer.

It will be appreciated that the present prostate cancer protein-modulating compounds can be administered alone or in combination with additional prostate cancer modulating compounds or with other therapeutic agent, e.g., other anti-cancer agents or treatments.

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In numerous embodiments, one or more nucleic acids, e.g., polynucleotides comprising nucleic acid sequences set forth in Tables 1-16, such as antisense polynucleotides or ribozymes, will be introduced into cells, in vitro or in vivo. The present invention provides methods, reagents, vectors, and cells useful for expression of prostate cancer-associated polypeptides and nucleic acids using in vitro (cell-free), ex vivo or in vivo (cell or organism-based) recombinant expression systems.

The particular procedure used to introduce the nucleic acids into a host cell for expression of a protein or nucleic acid is application specific. Many procedures for introducing foreign nucleotide sequences into host cells may be used. These include the use of calcium phosphate transfection, spheroplasts, electroporation, liposomes, microinjection, plasma vectors, viral vectors and any of the other well known methods for introducing cloned genomic DNA, cDNA, synthetic DNA or other foreign genetic material into a host cell (see, e.g., Berger & Kimmel, Guide to Molecular Cloning Techniques, Methods in Enzymology volume 152 (Berger), Ausubel et al., eds., Current Protocols (supplemented through 1999), and Sambrook et al., Molecular Cloning - A Laboratory Manual (2nd ed., Vol. 1-3, 1989.

In a preferred embodiment, prostate cancer proteins and modulators are administered as therapeutic agents, and can be formulated as outlined above. Similarly, prostate cancer genes (including both the full-length sequence, partial sequences, or regulatory sequences of the prostate cancer coding regions) can be administered in a gene therapy application. These prostate cancer genes can include antisense applications, either as gene therapy (i.e. for incorporation into the genome) or as antisense compositions, as will be appreciated by those in the art.

Prostate cancer polypeptides and polynucleotides can also be administered as vaccine compositions to stimulate HTL, CTL and antibody responses.. Such vaccine

compositions can include, e.g., lipidated peptides (see, e.g., Vitiello, A. et al., J. Clin. Invest. 95:341 (1995)), peptide compositions encapsulated in poly(DL-lactide-co-glycolide) ("PLG") microspheres (see, e.g., Eldridge, et al., Molec. Immunol. 28:287-294, (1991); Alonso et al., Vaccine 12:299-306 (1994); Jones et al., Vaccine 13:675-681 (1995)), peptide compositions contained in immune stimulating complexes (ISCOMS) (see, e.g., Takahashi et al., Nature 344:873-875 (1990); Hu et al., Clin Exp Immunol. 113:235-243 (1998)), multiple antigen peptide systems (MAPs) (see, e.g., Tam, Proc. Natl. Acad. Sci. U.S.A. 85:5409-5413 (1988); Tam, J. Immunol. Methods 196:17-32 (1996)), peptides formulated as multivalent peptides; peptides for use in ballistic delivery systems, typically crystallized peptides, viral delivery vectors (Perkus, et al., In: Concepts in vaccine development (Kaufmann, ed., p. 379, 1996); Chakrabarti, et al., Nature 320:535 (1986); Hu et al., Nature 320:537 (1986); Kieny, et al., AIDS Bio/Technology 4:790 (1986); Top et al., J. Infect. Dis. 124:148 (1971); Chanda et al., Virology 175:535 (1990)), particles of viral or synthetic origin (see, e.g., Kofler et al., J. Immunol. Methods. 192:25 (1996); Eldridge et al., Sem. Hematol. 30:16 (1993); Falo et al., Nature Med. 7:649 (1995)), adjuvants (Warren et al., Annu. Rev. Immunol. 4:369 (1986); Gupta et al., Vaccine 11:293 (1993)), liposomes (Reddy et al., J. Immunol. 148:1585 (1992); Rock, Immunol. Today 17:131 (1996)), or, naked or particle absorbed cDNA (Ulmer, et al., Science 259:1745 (1993); Robinson et al., Vaccine 11:957 (1993); Shiver et al., In: Concepts in vaccine development (Kaufmann, ed., p. 423, 1996); Cease & Berzofsky, Annu. Rev. Immunol. 12:923 (1994) and Eldridge et al., Sem. Hematol. 30:16 (1993)). Toxin-targeted delivery technologies, also known as receptor mediated targeting, such as those of Avant Immunotherapeutics, Inc. (Needham, Massachusetts) may also be used.

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Vaccine compositions often include adjuvants. Many adjuvants contain a substance designed to protect the antigen from rapid catabolism, such as aluminum hydroxide or mineral oil, and a stimulator of immune responses, such as lipid A, *Bortadella pertussis* or *Mycobacterium tuberculosis* derived proteins. Certain adjuvants are commercially available as, e.g., Freund's Incomplete Adjuvant and Complete Adjuvant (Difco Laboratories, Detroit, MI); Merck Adjuvant 65 (Merck and Company, Inc., Rahway, NJ); AS-2 (SmithKline Beecham, Philadelphia, PA); aluminum salts such as aluminum hydroxide gel (alum) or aluminum phosphate; salts of calcium, iron or zinc; an insoluble suspension of acylated tyrosine; acylated sugars; cationically or anionically derivatized polysaccharides;

polyphosphazenes; biodegradable microspheres; monophosphoryl lipid A and quil A. Cytokines, such as GM-CSF, interleukin-2, -7, -12, and other like growth factors, may also be used as adjuvants.

Vaccines can be administered as nucleic acid compositions wherein DNA or RNA encoding one or more of the polypeptides, or a fragment thereof, is administered to a patient. This approach is described, for instance, in Wolff *et. al.*, *Science* 247:1465 (1990) as well as U.S. Patent Nos. 5,580,859; 5,589,466; 5,804,566; 5,739,118; 5,736,524; 5,679,647; WO 98/04720; and in more detail below. Examples of DNA-based delivery technologies include "naked DNA", facilitated (bupivicaine, polymers, peptide-mediated) delivery, cationic lipid complexes, and particle-mediated ("gene gun") or pressure-mediated delivery (*see, e.g.*, U.S. Patent No. 5,922,687).

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For therapeutic or prophylactic immunization purposes, the peptides of the invention can be expressed by viral or bacterial vectors. Examples of expression vectors include attenuated viral hosts, such as vaccinia or fowlpox. This approach involves the use of vaccinia virus, e.g., as a vector to express nucleotide sequences that encode prostate cancer polypeptides or polypeptide fragments. Upon introduction into a host, the recombinant vaccinia virus expresses the immunogenic peptide, and thereby elicits an immune response. Vaccinia vectors and methods useful in immunization protocols are described in, e.g., U.S. Patent No. 4,722,848. Another vector is BCG (Bacille Calmette Guerin). BCG vectors are described in Stover et al., Nature 351:456-460 (1991). A wide variety of other vectors useful for therapeutic administration or immunization e.g. adeno and adeno-associated virus vectors, retroviral vectors, Salmonella typhi vectors, detoxified anthrax toxin vectors, and the like, will be apparent to those skilled in the art from the description herein (see, e.g., Shata et al., Mol Med Today 6:66-71 (2000); Shedlock et al., J Leukoc Biol 68:793-806 (2000); Hipp et al., In Vivo 14:571-85 (2000)).

Methods for the use of genes as DNA vaccines are well known, and include placing a prostate cancer gene or portion of a prostate cancer gene under the control of a regulatable promoter or a tissue-specific promoter for expression in a prostate cancer patient. The prostate cancer gene used for DNA vaccines can encode full-length prostate cancer proteins, but more preferably encodes portions of the prostate cancer proteins including peptides derived from the prostate cancer protein. In one embodiment, a patient is

immunized with a DNA vaccine comprising a plurality of nucleotide sequences derived from a prostate cancer gene. For example, prostate cancer-associated genes or sequence encoding subfragments of a prostate cancer protein are introduced into expression vectors and tested for their immunogenicity in the context of Class I MHC and an ability to generate cytotoxic T cell responses. This procedure provides for production of cytotoxic T cell responses against cells which present antigen, including intracellular epitopes.

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In a preferred embodiment, the DNA vaccines include a gene encoding an adjuvant molecule with the DNA vaccine. Such adjuvant molecules include cytokines that increase the immunogenic response to the prostate cancer polypeptide encoded by the DNA vaccine. Additional or alternative adjuvants are available.

In another preferred embodiment prostate cancer genes find use in generating animal models of prostate cancer. When the prostate cancer gene identified is repressed or diminished in cancer tissue, gene therapy technology, e.g., wherein antisense RNA directed to the prostate cancer gene will also diminish or repress expression of the gene. Animal models of prostate cancer find use in screening for modulators of a prostate cancer-associated sequence or modulators of prostate cancer. Similarly, transgenic animal technology including gene knockout technology, e.g. as a result of homologous recombination with an appropriate gene targeting vector, will result in the absence or increased expression of the prostate cancer protein. When desired, tissue-specific expression or knockout of the prostate cancer protein may be necessary.

It is also possible that the prostate cancer protein is overexpressed in prostate cancer. As such, transgenic animals can be generated that overexpress the prostate cancer protein. Depending on the desired expression level, promoters of various strengths can be employed to express the transgene. Also, the number of copies of the integrated transgene can be determined and compared for a determination of the expression level of the transgene. Animals generated by such methods find use as animal models of prostate cancer and are additionally useful in screening for modulators to treat prostate cancer.

Kits for Use in Diagnostic and/or Prognostic Applications

For use in diagnostic, research, and therapeutic applications suggested above, kits are also provided by the invention. In the diagnostic and research applications such kits

may include any or all of the following: assay reagents, buffers, prostate cancer-specific nucleic acids or antibodies, hybridization probes and/or primers, antisense polynucleotides, ribozymes, dominant negative prostate cancer polypeptides or polynucleotides, small molecules inhibitors of prostate cancer-associated sequences *etc*. A therapeutic product may include sterile saline or another pharmaceutically acceptable emulsion and suspension base.

In addition, the kits may include instructional materials containing directions (i.e., protocols) for the practice of the methods of this invention. While the instructional materials typically comprise written or printed materials they are not limited to such. Any medium capable of storing such instructions and communicating them to an end user is contemplated by this invention. Such media include, but are not limited to electronic storage media (e.g., magnetic discs, tapes, cartridges, chips), optical media (e.g., CD ROM), and the like. Such media may include addresses to internet sites that provide such instructional materials.

The present invention also provides for kits for screening for modulators of prostate cancer-associated sequences. Such kits can be prepared from readily available materials and reagents. For example, such kits can comprise one or more of the following materials: a prostate cancer-associated polypeptide or polynucleotide, reaction tubes, and instructions for testing prostate cancer-associated activity. Optionally, the kit contains biologically active prostate cancer protein. A wide variety of kits and components can be prepared according to the present invention, depending upon the intended user of the kit and the particular needs of the user. Diagnosis would typically involve evaluation of a plurality of genes or products. The genes will be selected based on correlations with important parameters in disease which may be identified in historical or outcome data.

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EXAMPLES

Example 1: Tissue Preparation, Labeling Chips, and Fingerprints

5 Purifying total RNA from tissue sample using TRIzol Reagent

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The sample weight is first estimated. The tissue samples are homogenized in 1 ml of TRIzol per 50 mg of tissue using a homogenizer (e.g., Polytron 3100). The size of the generator/probe used depends upon the sample amount. A generator that is too large for the amount of tissue to be homogenized will cause a loss of sample and lower RNA yield. A larger generator (e.g., 20 mm) is suitable for tissue samples weighing more than 0.6 g. Fill tubes should not be overfilled. If the working volume is greater than 2 ml and no greater than 10 ml, a 15 ml polypropylene tube (Falcon 2059) is suitable for homogenization.

Tissues should be kept frozen until homogenized. The TRIzol is added directly to the frozen tissue before homogenization. Following homogenization, the insoluble material is removed from the homogenate by centrifugation at 7500 x g for 15 min. in a Sorvall superspeed or 12,000 x g for 10 min. in an Eppendorf centrifuge at 4°C. The cleared homogenate is then transferred to a new tube(s). Samples may be frozen and stored at -60 to -70°C for at least one month or else continue with the purification.

The next process is phase separation. The homogenized samples are incubated for 5 minutes at room temperature. Then, 0.2 ml of chloroform per 1ml of TRIzol reagent is added to the homogenization mixture. The tubes are securely capped and shaken vigorously by hand (do not vortex) for 15 seconds. The samples are then incubated at room temp. for 2-3 minutes and next centrifuged at 6500 rpm in a Sorvall superspeed for 30 min. at 4oC.

The next process is RNA Precipitation. The aqueous phase is transferred to a fresh tube: The organic phase can be saved if isolation of DNA or protein is desired. Then 0.5 ml of isopropyl alcohol is added per 1ml of TRIzol reagent used in the original homogenization. Then, the tubes are securely capped and inverted to mix. The samples are then incubated at room temp. for 10 minutes an centrifuged at 6500 rpm in Sorvall for 20 min. at 4°C.

The RNA is then washed. The supernatant is poured off and the pellet washed with cold 75% ethanol. 1 ml of 75% ethanol is used per 1 ml of the TRIzol reagent used in the initial homogenization. The tubes are capped securely and inverted several times to loosen pellet without vortexing. They are next centrifuged at $<8000\,$ rpm ($<7500\,$ x g) for 5 minutes at 4° C.

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The RNA wash is decanted. The pellet is carefully transferred to an Eppendorf tube (sliding down the tube into the new tube by use of a pipet tip to help guide it in if necessary). Tube(s) sizes for precipitating the RNA depending on the working volumes. Larger tubes may take too long to dry. Dry pellet. The RNA is then resuspended in an appropriate volume (e.g., 2 -5 ug/ul) of DEPC H₂0. The absorbance is then measured.

The poly A+ mRNA may next be purified from total RNA by other methods such as Qiagen's RNeasy kit. The poly A + mRNA is purified from total RNA by adding the oligotex suspension which has been heated to 37°C and mixing prior to adding to RNA. The Elution Buffer is incubated at 70°C. If there is precipitate in the buffer, warm up the 2 x Binding Buffer at 65°C. The the total RNA is mixed with DEPC-treated water, 2 x Binding Buffer, and Oligotex according to Table 2 on page 16 of the Oligotex Handbook and next incubated for 3 minutes at 65°C and 10 minutes at room temperature.

The preparation is centrifuged for 2 minutes at 14,000 to 18,000 g, preferably, at a "soft setting," The supernatant is removed without disturbing Oligotex pellet. A little bit of solution can be left behind to reduce the loss of Oligotex. The supernatant is saved until satisfactory binding and elution of poly A+ mRNA has been found.

Then, the preparation is gently resuspended in Wash Buffer OW2 and pipetted onto the spin column and centrifuged at full speed (soft setting if possible) for 1 minute.

Next, the spin column is transferred to a new collection tube and gently resuspended in Wash Buffer OW2 and centrifuged as described herein.

Then, the spin column is transferred to a new tube and eluted with 20 to 100 ul of preheated (70°C) Elution Buffer. The Oligotex resin is gently resuspended by pipetting up and down. The centrifugation is repeated as above and the elution repeated with fresh elution buffer or first eluate to keep the elution volume low.

The absorbance is next read to determine the yield, using diluted Elution Buffer as the blank.

Before proceeding with cDNA synthesis, the mRNA is precipitated before proceeding with cDNA synthesis, as components leftover or in the Elution Buffer from the Oligotex purification procedure will inhibit downstream enzymatic reactions of the mRNA. 0.4 vol. of 7.5 M NH4OAc + 2.5 vol. of cold 100% ethanol is added and the preparation precipitated at -20°C 1 hour to overnight (or 20-30 min. at -70°C), and centrifuged at 14,000-16,000 x g for 30 minutes at 4°C. Next, the pellet is washed with 0.5 ml of 80% ethanol (-20°C) and then centrifuged at 14,000-16,000 x g for 5 minutes at room temperature. The 80% ethanol wash is then repeated. The last bit of ethanol from the pellet is then dried without use of a speed vacuum and the pellet is then resuspended in DEPC H₂0 at 1ug/ul concentration.

Alternatively the RNA may be purified using other methods (e.g., Qiagen's RNeasy kit).

No more than 100 ug is added to the RNeasy column. The sample volume is adjusted to 100 ul with RNase-free water. 350 ul Buffer RLT and then 250 ul ethanol (100%) are added to the sample. The preparation is then mixed by pipetting and applied to an RNeasy mini spin column for centrifugation (15 sec at >10,000 rpm). If yield is low, reapply the flowthrough to the column and centrifuge again.

Then, transfer column to a new 2 ml collection tube and add 500 ul Buffer RPE and centrifuge for 15 sec at >10,000 rpm. The flowthrough is discarded. 500 ul Buffer RPE and is then added and the preparation is centriuged for 15 sec at >10,000 rpm. The flowthrough is discarded. and the column membrane dried by centrifuging for 2 min at maximum speed. The column is transferred to a new 1.5-ml collection tube. 30-50 ul of RNase-free water is applied directly onto column membrane. The column is then centrifuged for 1 min at >10,000 rpm and the elution step repeated.

The absorbance is then read to determine yield. If necessary, the material may be ethanol precipitated with ammonium acetate and 2.5X volume 100% ethanol.

First Strand cDNA Synthesis

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The first strand can be make using using Gibco's "SuperScript Choice System for cDNA Synthesis" kit. The starting material is 5 ug of total RNA or 1 ug of polyA+ mRNAl. For total RNA, 2 ul of SuperScript RT is used; for polyA+ mRNA, 1 ul of SuperScript RT is used. The final volume of first strand synthesis mix is 20 ul. The RNA should be in a volume no greater than 10 ul. The RNA is incubated with 1 ul of 100 pmol T7-T24 oligo for 10 min at 70°C followed by addition on ice of 7 ul of: 4ul 5X 1st Strand Buffer, 2 ul of 0.1M DTT, and 1 ul of 10mM dNTP mix. The preparation is then incubated at 37°C for 2 min before addition of the SuperScript RT followed by incubation at 37°C for 1 hour.

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Second Strand Synthesis

For the second strand synthesis, place 1st strand reactions on ice and add: 91 ul DEPC H₂0; 30 ul 5X 2nd Strand Buffer; 3 ul 10mM dNTP mix; 1 ul 10 U/ul E.coli DNA Ligase; 4 ul 10 U/ul E.coli DNA Polymerase; and 1 ul 2 U/ul RNase H. Mix and incubate 2 hours at 16°C. Add 2 ul T4 DNA Polymerase. Incubate 5 min at 16°C. Add 10 ul of 0.5M EDTA.

Cleaning up cDNA

The cDNA is purified using Phenol:Chloroform:Isoamyl Alcohol (25:24:1) and Phase-Lock gel tubes. The PLG tubes are centrifuged for 30 sec at maximum speed. The cDNA mix is then transferred to PLG tube. An equal volume of phenol:chloroform:isamyl alcohol is then added, the preparation shaken vigorously (no vortexing), and centrifuged for 5 minutes at maximum speed. The top aqueous solution is transferred to a new tube and ethanol precipitated by adding 7.5X 5M NH4OAc and 2.5X volume of 100% ethanol. Next, it is centrifuged immediately at room temperature for 20 min, maximum speed. The supernatant is removed, and the pellet washed with 2X with cold 80% ethanol. As much ethanol wash as possible should be removed before air drying the pellet; and resuspending it in 3 ul RNase-free water.

In vitro Transcription (IVT) and labeling with biotin

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In vitro Transcription (IVT) and labeling with biotin is performed as follows: Pipet 1.5 ul of cDNA into a thin-wall PCR tube. Make NTP labeling mix by combining 2 ul T7 10xATP (75 mM) (Ambion); 2 ul T7 10xGTP (75 mM) (Ambion); 1.5 ul T7 10xCTP (75 mM) (Ambion); 1.5 ul T7 10xUTP (75 mM) (Ambion); 3.75 ul 10 mM Bio-11-UTP (Boehringer-Mannheim/Roche or Enzo); 3.75 ul 10 mM Bio-16-CTP (Enzo); 2 ul 10x T7 transcription buffer (Ambion); and 2 ul 10x T7 enzyme mix (Ambion). The final volume is 20 ul. Incubate 6 hours at 37°C in a PCR machine. The RNA can be furthered cleaned. Clean-up follows the previous instructions for RNeasy columns or Qiagen's RNeasy protocol handbook. The cRNA often needs to be ethanol precipitated by resuspension in a volume compatible with the fragmentation step.

Fragmentation is performed as follows. 15 ug of labeled RNA is usually fragmented. Try to minimize the fragmentation reaction volume; a 10 ul volume is recommended but 20 ul is all right. Do not go higher than 20 ul because the magnesium in the fragmentation buffer contributes to precipitation in the hybridization buffer. Fragment RNA by incubation at 94 C for 35 minutes in 1 x Fragmentation buffer (5 x Fragmentation buffer is 200 mM Tris-acetate, pH 8.1; 500 mM KOAc; 150 mM MgOAc). The labeled RNA transcript can be analyzed before and after fragmentation. Samples can be heated to 65°C for 15 minutes and electrophoresed on 1% agarose/TBE gels to get an approximate idea of the transcript size range.

For hybridization, 200 ul (10 ug cRNA) of a hybridization mix is put on the chip. If multiple hybridizations are to be done (such as cycling through a 5 chip set), then it is recommended that an initial hybridization mix of 300 ul or more be made. The hybridization mix is: fragment labeled RNA (50 ng/ul final conc.); 50 pM 948-b control oligo; 1.5 pM BioB; 5 pM BioC; 25 pM BioD; 100 pM CRE; 0.1 mg/ml herring sperm DNA; 0.5 mg/ml acetylated BSA; and 300 ul with 1xMES hyb buffer.

The hybridization reaction is conducted with non-biotinylated IVT (purified by RNeasy columns) (see example 1 for steps from tissue to IVT): The following mixture is prepared:

IVT antisense RNA; 4 μ g: μ l Random Hexamers (1 μ g/ μ l): 4 μ l H₂O: μ l 14 μ l

5 Incubate the above 14 µl mixture at 70°C for 10 min.; then put on ice.

The Reverse transcription procedure uses the following mixture:

The above solution is added to the hybridization reaction and incubated for 30 min., 42°C.

Then, 1 µl SSII is added and incubated for another hour before being placed on ice.

The 50X dNTP mix contains 25mM of cold dATP, dCTP, and dGTP, 10mM of dTTP and is made by adding 25 μl each of 100mM dATP, dCTP, and dGTP; 10 μl of 100mM dTTP to 15 μl H₂O.]

RNA degradation is performed as follows. Add 86 µl H2O, 1.5 µl 1M NaOH/2 mM EDTA and incubate at 65°C, 10 min.. For U-Con 30, 500 µl TE/sample spin at 7000 g for 10 min, save flow through for purification. For Qiagen purification, suspend u-con recovered material in 500 µl buffer PB and proceed using Qiagen protocol. For DNAse digestion, add 1 ul of 1/100 dilution of DNAse/30 ul Rx and incubate at 37°C for 15 min. Incubate at 5 min 95°C to denature the DNAse.

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Sample preparation

For sample preparation, add Cot-1 DNA, 10 μ l; 50X dNTPs, 1 μ l; 20X SSC, 2.3 μ l; Na pyro phosphate, 7.5 μ l; 10 mg/ml Herring sperm DNA; 1 μ l of 1/10 dilution to 21.8 final vol. Dry in speed vac. Resuspend in 15 μ l H₂0. Add 0.38 μ l 10% SDS. Heat

95°C, 2 min and slow cool at room temp. for 20 min. Put on slide and hybridize overnight at 64°C. Washing after the hybridization: 3X SSC/0.03% SDS: 2 min., 37.5 ml 20X SSC+0.75ml 10% SDS in 250ml H₂O; 1X SSC: 5 min., 12.5 ml 20X SSC in 250ml H₂O; 0.2X SSC: 5 min., 2.5 ml 20X SSC in 250ml H₂O. Dry slides and scan at appropriate PMT's and channels.

Example 2: Taxol resistant Xenograft Model of Human Prostate Cancer

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Treatment regimens that include paclitaxel (Taxol; Bristol-Myers Squibb

Company, Princeton, NJ) have been particularly successful in treating hormone-refractory prostate cancer in the phase II setting (Smith et al., Semin. Oncol. 26(1 Suppl 2):109-11 (1999)). However, many patients develop tumors which are initially, or later become, resistant to taxol. To identify genes that may be involved with resistance to taxol, or are regulated in response to taxol resistance, and therefore may be used to treat, or identify, taxol resistance in patients, the following experiments were carried out.

The androgen-independent human cell line CWR22R was grown as a xenograft in nude mice (Nagabhushan et al., Cancer Res. 56(13):3042-3046 (1996); Agus et al., J. Natl. Cancer Inst.91(21):1869-1876 (1999); Bubendorf et al., J. Natl. Cancer Inst. 91(20):1758-1764 (1999)). Initially, these xenograft tumors were sensitive to therapeutic doses of taxol. The mice were treated continuously with sub-therapeutic doses, and the tumors were allowed to grow for 3-4 weeks, before surgical removal of the tumors. The tumor from an individual mouse was then minced, and a small portion was then injected into a healthy nude mouse, establishing a second

passage of the tumor. This mouse was then treated continuously with the same sub-therapeutic dose of taxol. This process was repeated 14 times, and a portion of each generation of xenograft tumor was collected. There was increasing resistance to therapeutic doses of taxol with each generation. Bythe end of the process, the tumors were fully resistant to therapeutic doses of taxol. RNA from each generation of tumor was then isolated, and individual mRNA species were quantified using a custom Affymetrix GeneChip® oligonucleotide microarray, with probes to interrogate approximately 35,000

unique mRNA transcripts. Genes were selected that showed a statistically significant up-regulation, or down-regulation, during the subsequent generations of increasingly taxol-resistant tumors. Only one gene was significantly up-regulated, whereas 24 genes were down-regulated; these are presented in Table 10.

The gene sequences identified to be overexpressed in prostate cancer may be used to identify coding regions from the public DNA database. The sequences may be used to either identify genes that encode known proteins, or they may be used to predict the coding regions from genomic DNA using exon prediction algorithms, such as FGENESH (Salamov and Solovyev, 2000, Genome Res. 10:516-522). In addition, one of ordinary skill in the art would understand how to obtain the unigene cluster identification and sequence information according to the exemplar accession numbers provided in Tables 1-16. (see, http://www.ncbi.nlm.nih.gov/UniGene/).

TABLE1: shows genes, including expression sequence tags, differentially expressed in prostate tumor tissue compared to normal tissue as analyzed using the Affymetrix/Eos Hu01
 GeneChip array. Shown are the relative amounts of each gene expressed in prostate tumor samples and various normal tissue samples showing the highest expression of the gene.

10	Pkey: ExAcon:	Unique Eos probeset identifier number Exemplar Accession number, Genbank accession number
10	UnigenelD:	Unigene number
	Unigene Title:	Unigene gene title
	R1:	Ratio of tumor to normal body tissue

				•	
15					
10	Pkey	UnigeneiD	ExAccn	Uningene Title	R1
	131919	Hs.272458	AA121266	ESTs .	37.2
		Hs.290905		ESTs; Weakly similar to (defline not ava	32.6
20		Hs.31412		ESTs	30.1
20		Hs.1852		acid phosphatase; prostate	25.2
		Hs.279477		ESTs	24.8
		Hs.183752		microseminoprotein; beta-	23.8
		Hs.171995		kallikrein 3; (prostate specific antigen	21.4
25		Hs.57771		Homo sapiens mRNA for serine protease (T	18.9
		Hs.162859		ESTs	18.6
		Hs.30343		ESTs	17.4
		Hs.1832	K01911	neuropeptide Y	17.3
		Hs.1915		folate hydrolase (prostate-specific memb	17
30		Hs.40808		ESTs	16.9
50		Hs.262476		S-adenosylmethionine decarboxylase 1	16.7
		Hs.33287		ESTs	16.5
		Hs.11260		ESTs	16.4
				Antigen, Prostate Specific, Alt. Splice	16
35		Hs.181350		kallikrein 2; prostatic	15.4
33		Hs.99872		fetal Alzheimer antigen	15
		Hs.62192		coagulation factor III (thromboplastin;	13.9
		Hs.8236	D62633	ESTs	12.7
		Hs.7780		ESTs	12.5
40		Hs.193380		ESTs .	12.3
		Hs.2178	X57985	H2B histone family; member Q	11.8
		Hs.182339		ESTs	11.8
		Hs.172129		yp19h1.r1 Soares breast 3NbHBst Homo sap	11.8
		Hs.102720		ESTs	11.6
45		Hs.14846		ESTs	11.4
		Hs.78045		actin; gamma 2; smooth muscle; enteric	11
		Hs.268744		ESTs; Moderately similar to KIAA0273 [H.	10.9
		126645	Al167942	Homo sapiens BAC done RG041D11 from 7q2	10.7
		Hs.95420	N40141	Homo sapiens mRNA for JM27 protein; comp	10.6
50		Hs.113314		ESTs	- 10.6
	118417		N66048	ESTs; Weakly similar to polymerase [H.sa	10.5
	126758	Hs.293960	W37145	ESTs	10.2
		Hs.8364		ESTs	10.1
			AA236324 -	ESTs; Weakly similar to !!!! ALU CLASS A	10.1
55		Hs.30652		ESTs	10.1
		Hs.15641		ESTs	10.1
	115719	Hs.59622	AA416997	ESTs	10
	123209	Hs.203270	AA489711	ESTs	9.9
	101664	Hs.121017	M60752	H2A histone family; member A	9.8
60	112971	Hs.83883	T17185	ESTs	9.7
-	102519	Hs.80296	U52969	Purkinje cell protein 4	9.7
	117984	Hs.106778	N51919 ·	ESTs	9.7
		Hs.22209		ESTs	9.4
	129523	Hs.274509	M30894	T-cell receptor; gamma cluster	9.4
65	132964	Hs.167133	AA031360	ESTs	9.2
	121853	Hs.98502	AA425887	ESTs	9

	115764	Hs.91011	AA421562	anterior gradient 2 (Xenopus laevis; sec	8.9
		Hs.55999	W47380	ESTs	8.9
	100552	Hs.301946	HG2167-HT2237	Protein Kinase Ht31, Camp-Dependent	8.9
_	105627	Hs.23317	AA281245	ESTs	8.8
5		Hs.76422	M22430	phospholipase A2; group IIA (platelets;	8.7
		Hs.31146	AA456264	ESTs; Highly similar to (defline not ava	8.5 8.5
		Hs.293185 Hs.49397	N67889	yz61c5.s1 Soares_multiple_sclerosis_2NbH ESTs	8.2
		Hs.76704	T68510	ESTs	8.2
10		Hs.334762		ESTs; Weakly similar to KIAA0319 [H.sapi	8.1
•		Hs.20415	AA402000	ESTs; Weakly similar to GS3786 [H.sapien	8
		Hs.278695	N95796	ESTs	8
	100394	Hs.66052	D84276	CD38 antigen (p45)	8
		Hs.24192	Z38688	ESTs	7.9
15		Hs.301527		tumor necrosis factor (ligand) superfami	7.7
		Hs.23023	AA456135	ESTs	7.6 7.5
		Hs.105700 Hs.72472	AA250737	secreted frizzled-related protein 4 ESTs	7.4
		Hs.22627	R43162	ESTs	7.1
20	102398	110122021	U42359	Human N33 protein form 1 (N33) gene, exo	7
		Hs.2256	L22524	matrix metalloproteinase 7 (matrilysin;	6.9
		Hs.288462	AA195718	ESTs	6.9
		Hs.169849		myosin-binding protein C; slow-type	6.9
25		Hs.155691		pre-B-cell leukemia transcription factor	6.8
25		Hs.302267 Hs.257924		ESTs; Weakly similar to W01A6.c [C.elega ESTs	6.8 6.8
		Hs.326416		ESTs	6.7
-		Hs.173684		ESTs; Weakly similar to (defline not ava	6.7
		Hs.171995		kallikrein 3; (prostate specific antigen	6.6
30		Hs.26691	AA219134	ESTs	6.6
		Hs.16193		Homo sapiens mRNA; cDNA DKFZp586B211 (fr	6.6
		Hs.59838	AA490969	ESTs	6.6
		Hs.323378		H.sapiens mRNA for transmembrane protein	6.6 6.5
35		Hs.75746 Hs.278628	U07919	aldehyde dehydrogenase 6 ESTs; Moderately similar to APXL gene pr	6.5
55		Hs.108787		Homo sapiens Mcd4p homolog mRNA; complet	6.5
		Hs.126085		ESTs	6.5
		Hs.3383	AA010163	upstream regulatory element binding prot	6.5
40		Hs.7232	T23670	ESTs	6.4
40		Hs.8768	AA236559	ESTs; Weakly similar to neuronal thread	6.4
		Hs.26289	AA009527	ESTS	6.4 6.3
		Hs.334786 Hs.15113	AF000573	Human HF.12 gene mRNA homogentisate 1;2-dioxygenase (homogenti	6.3
		Hs.278428		Homo sapiens mRNA for KIAA0896 protein;	6.3
45		Hs.250528		ESTs; Weakly similar to ANKYRIN; BRAIN V	6.3
	133061	Hs.296638	AB000584	prostate differentiation factor	6.3
		Hs.279923		ESTs; Weakly similar to similar to GTP-b	6.2
		Hs.878	L29008	sorbitol dehydrogenase	6.2
50		Hs.37744		ESTs EST27953 Cerebellum II Homo sapiens cDNA	6.2 6.2
.50	127248	Hs.179902	AA325029 H04108	ESTs; Weakly similar to (define not ava	6.2
	105500	Hs.222399	AA256485	ESTs .	6.1
		Hs.2714	X74142	forkhead (Drosophila)-like 1 -	6.1
		Hs.40289		ESTs	6
55		Hs.203213		ESTs	5.9
		Hs.72988	AA281793	ESTs	5.8
		Hs.301997		ESTs	5.7 5.7
		Hs.48948	AA491457 AA034020	ESTs ESTs	5.7
60		Hs.61539 Hs.125019		ESTS; Weakly similar to !!!! ALU SUBFAMI	5.6
J	106375	Hs.289072	AA443993	ESTs	5.6
		Hs.170195		bone morphogenetic protein 7 (osteogenic	5.6
		Hs.140237	_	ESTs; Weakly similar to neuronal thread	5.6
~~		Hs.337616		phosphodiesterase 3B; cGMP-inhibited	5.6
65		Hs.62354		Human beige-like protein (BGL) mRNA; par	5.5 5.5
		Hs.45107 Hs.281434		ESTs heat shock 70kD protein 1	5.5 5.5
		Hs.98732		Homo sapiens Chromosome 16 BAC clone CIT	5.5
		Hs.262476		S-adenosylmethionine decarboxylase 1	5.5

			1110100		
	113938	0.10045	W81598	ESTs	5.4
		Hs.246315		ESTs	5.4
•		Hs.75722	A1283493	ribophorin II	5.4
F		Hs.80120		UDP-N-acetyl-alpha-D-galactosamine:polyp	5.4
5		Hs.7780	AA056482	ESTs	5.3
		Hs.21223		calponin 1; basic; smooth muscle	5.3
		Hs.326392		Human guanine nucleotide exchange factor	5.3
		Hs.98944	AA365031	ESTs	5.3
10		Hs.167531		ESTs; Weakly similar to (defline not ava	5.3
10		Hs.108336		ESTs; Weakly similar to IIII ALU SUBFAMI	5.3
		Hs.25351	U90304	iroquois-class homeodomain protein	5.3
		Hs.194369		Homo sapiens chromosome 1 atrophin-1 rel	5.3
		Hs.109201		ESTs; Highly similar to (defline not ava	5.2 5.2
15		Hs.79428 Hs.159872	U15174	BCL2/adenovirus E1B 19kD-interacting pro ESTs	5.2
13	10/240	TIS. 10001Z	AA027317	ESTs; Weakly similar to !!!! ALU SUBFAMI	5.2
		Hs.108327		damage-specific DNA binding protein 1 (1	5.2
		Hs.194228		ESTs; Moderately similar to IIII ALU SUB	5.2
		Hs.195850		keratin 5 (epidermolysis bullosa simplex	5.1
20		Hs.184598		ESTs; Weakly similar to !!!! ALU SUBFAMI	5.1
20		Hs.281428		ESTs; Moderately similar to !!!! ALU SUB	5.1
		Hs.169119		ESTs	5.1
		Hs.54416	X91868	sine oculis homeobox (Drosophila) homolo	5.1
		Hs.106778		ESTs; Highly similar to (defline not ava	5.1
25		Hs.148932		ESTs; Moderately similar to semaphorin V	5.1
		Hs.226434		ESTs	5.1
	116238	Hs.47144	AA479362	ESTs	5
	102913	Hs.80342	X07696	keratin 15	5
	103011	Hs.326035		early growth response 1	5
30	126023		H58881	yr36d09.r1 Soares fetal liver spleen 1NF	5
		Hs.13804	AA037316	ESTs	5
		Hs.39288		ESTs; Weakly similar to !!!! ALU SUBFAMI	5
		Hs.89732	X78932	zinc finger protein 273	5
35		Hs.23311	AB002365	Human mRNA for KIAA0367 gene; partial cd	4.9
33		Hs.3796	D83492	EphB6	4.9 4.9
		Hs.177537	F10523	ESTs primase; polypeptide 2A (58kD)	4.9
		Hs.74519 Hs.71119	U42360	Human N33 mRNA; complete cds	4.8
	104776	113.71113	AA026349	ESTs	4.8
40		Hs.128749		Homo sapiens alpha-methylacyl-CoA racema	4.8
		Hs.143087		ESTs	4.8
		Hs.26009	W86307	Homo sapiens mRNA for KIAA0860 protein;	4.8
	105288	Hs.3585	AA233168	ESTs; Weakly similar to coded for by C.	4.8
	135035	Hs.284186	H89575	ESTs	4.8
45	104144	Hs.183390	AA447439	ESTs; Weakly similar to ZINC FINGER PROT	4.8
		Hs.288126	AA621604	ESTs	4.8
	125982		R98091	RAE1 (RNA export 1; S.pombe) homolog	4.8
		Hs.26243		ESTs	4.8
50		Hs.117950		multifunctional polypeptide similar to S	4.7
50	129735	Na 400000	W80701	ESTs; Weakly similar to HERV-E envelope	4.7
		Hs.106390		ESTs	4.7 4.7
	103731	Hs.127602	AA070545	zm7c3.r1 Stratagene neuroepithelium (#93 ESTs	4.7
		Hs.231500		Human alucose transporter-like protein-l	4.7
55		Hs.1674	M90516	glutamine-fructose-6-phosphate transamin	4.7
		Hs.91622	H06373	Homo sapiens clone 24456 mRNA sequence	4.7
		Hs.82007	D42084	Human mRNA for KIAA0094 gene; partial cd	4.7
		Hs.89576	A1479264	ESTs	4.7
	131836	Hs.32990	AA610086	ESTs	4.7
60		Hs.239489	AA465093	TIA1 cytotoxic granule-associated RNA-bi	4.7
	114542	Hs.91011	AA055768	ESTs	4.6
	103806		AA130614	zo1f2.r1 Stratagene neuroepithelium NT2R	4.6
	130529		AA173238	small inducible cytokine A5 (RANTES)	4.6
(=		Hs.82065	AA406546	ESTs	4.6
65		Hs.293798		ESTs	4.6
		Hs.29679	AA452411	ESTs	4.6
		Hs.14158 Hs.100070	W86835	copine III	4.6
	10077/	Hs.89603	M91493 HG371-HT1063	EST Mucin 1, Epithelial, Alt. Splice 6	4.6 4.6
	100774		110071-1111003	move of charamit at object a	7.0

				Ret Transforming Gene	4.6
		Hs.3731	D11900	ESTs	4.6
	126086		H70975	yr73g01.r1 Soares fetal liver spleen 1NF	4.6
_		Hs.173094		ESTs	4.6
5		Hs.20166		Prostate stem cell antigen	4.6
	126959	11-00447	AA199853	ESTs; Moderately similar to IIII ALU SUB	4.5 4.5
		Hs.29117		H.sapiens mRNA for pur alpha extended 3'	4.5 4.5
		Hs.20953	AA039481	ESTS	4.5
10	125661	U- 004700	R50319	ESTs	4.5
10		Hs.234726		alpha-1-antichymotrypsin ESTs	4.5
		Hs.199160	X06272	signal recognition particle receptor ('d	4.5
		Hs.75730		Small Nuclear Ribonucleoprotein U1, 1snr	4.5
		Hs.7956	AA425906	ESTs	4.5
15		Hs.317584		ESTs	4.5
13		Hs.24758		ESTs	4.5
		Hs.44566		Human protein immuno-reactive with anti-	4.4
•		Hs.38176		Homo saplens mRNA for KIAA0606 protein;	4.4
		Hs.198760		neurofilament; heavy potypeptide (200kD)	4.4
20		Hs.1846	M22898	tumor protein p53 (Li-Fraumeni syndrome)	4.4
20		Hs.284296		ESTs; Highly similar to surface 4 integr	4.4
		Hs.22514		ESTs	4.4
		Hs.119394		ESTs	4.4
		Hs.29894		ESTs	4.4
25	121944	Hs.98518	AA429278	ESTs	4.4
	134401	Hs.211577	AA243746	ESTs; Highly similar to CG1 protein [H.s	4.4
	126458	Hs.288969	AA815252	ESTs; Weakly similar to !!!! ALU SUBFAMI	4.4
	133435	Hs.323966	T23983	ESTs; Moderately similar to !!!! ALU SUB	4.4
	105178	Hs.21941	AA187490	ESTs	4.3
30	127315		AA840834	nr27b06.r1 NCI_CGAP_Pr3 Homo sapiens cDN	4.3
		Hs.54424		H.saplens mRNA for hepatocyte nuclear fa	4.3
		Hs.282990		ESTs; Weakly similar to F52C12.2 [C.eleg	4.3
		Hs.47567	N52876	EST	4.3
25		Hs.278427		cerebellar degeneration-related protein	4.3
35		Hs.114688		ESTs	4.3 4.3
		Hs.105130		EST ESTs	4.3
		Hs.184245		ESTS	4.3
	109175	Hs.173540	AA180496	ESTs; Weakly similar to (defline not ava	4.3
40		Hs.46638	U57911	chromosome 11 open reading trame 8	4.3
40		Hs.79993	U88871	peroxisomal biogenesis factor 7	4.3
		Hs.5462	AF007216	solute carrier family 4; sodium bicarbon	4.3
		Hs.18946	AA094720	ESTs; Weakly similar to (defline not ava	4.3
		Hs.295923		seven in absentia (Drosophila) homolog 1	4.3
45		Hs.93872		ESTs	4.3
		Hs.334762		ESTs; Wealdy similar to KIAA0319 [H.sapi	4.2
		Hs.98747	AA431732	EST	4.2
	133136	Hs.6574	AF007165	suppressin (nuclear deformed epidermal a	4.2
	109890	Hs.20843	H04649	ESTs	4.2
50	133294	Hs.69997	R79723	H.sapiens mRNA for translin associated z	4.2
		Hs.83190	S80437	fatty acid synthase (3' region) [human,	4.2
		Hs.251064		NBR2	4.2
		Hs.27413		ESTs -	4.2
		Hs.248210		H.sapiens Mahlavu hepatocellular carcino	4.2
55		Hs.59815	W99362	EST	4.2
		Hs.283978		ESTs; Highly similar to (defline not ava	4.2
		Hs.1179		TATA box binding protein (TBP)-associate	4.2 4.2
	106566		AA455921	ESTs; Weakly similar to IIII ALU SUBFAMI	4.2
<i>c</i> 0		Hs.29852		ESTS	4.2
60		Hs.279929		H.sapiens mRNA for gp25L2 protein transcriptional repressor	4.2
		Hs.57419		ESTs	4.2
		Hs.326292			4.2
		Hs.94109		ESTs lactotransferrin	4.1
65		Hs.105938		yg95c6.r1 Soares infant brain 1NIB Homo	4.1
UJ		Hs.108850 Hs.6641	N98707	kinesin family member 5C	4.1
		Hs.14051		ESTs	4.1
		Hs.45032		ESTs	4.1
		Hs.327179		solute carrier family 17 (sodium phospha	4.1
	101010	. 10.067 175		and a farming the farming bearing	

		Hs.100861		ESTs; Weakly similar to p60 katanin [H.s	4.1
	130555	Hs.116774	AA450324	ESTs	4.1
	105765	Hs.24183	AA343514	ESTs	4.1
		Hs.26369	AA133237	ESTs	4.1
5				ESTs	4.1
J		Hs.181889			
		Hs.172129		ESTs; Moderately similar to IIII ALU SUB	4.1
	100234	Hs.3085	D29677	KIAA0054 gene product	4.1
	100959	Hs.118127	J00073	actin; alpha; cardiac muscle	4.1
		Hs.12913	AA620582	ESTs; Weakly similar to (defline not ava	4.1
10			AA128486	ESTs	4.1
10		Hs.8859			
		Hs.226795		glutathione S-transferase pi	4.1
	113056	Hs.8036	T26471	ESTs; Moderately similar to !!!! ALU SUB	4
	102460	Hs.211582	U48959	Homo sapiens myosin light chain kinase (4
		Hs.26813		ESTs; Weakly similar to (defline not ava	4
15		Hs.104207		ESTs (Common to Common to	4
IJ					4
		Hs.267967		ESTs; Weakly similar to !!!! ALU SUBFAMI	
	105329	Hs.22862	AA234561	ESTs	4
	115504	Hs.42736	AA291946	ESTs	4
	120726	Hs.97293	AA293656 .	ESTs	4
20		Hs.94560	Z26317	desmoglein 2	4
20				ESTS	4
		Hs.144941			
	106394	Hs.25320	AA447223	ESTs	4
	128046		AA873285	ESTs	4
	103391	Hs.114366	X94453	pyrroline-5-carboxylate synthetase (glut	4
25		Hs.27004	AA449455	ESTs	4
2,5					4
		Hs.86276	W27601	ESTs; Moderately similar to (defline not	
	129593	Hs.98314	AA487015	ESTs; Weakly similar to !!!! ALU SUBFAMI	3.9
	110151	Hs.31608	H18836	ESTs ·	3.9
	105344	Hs.8645	AA235303	ESTs	3.9
30		Hs.301871		ESTs	3.9
50				ESTs	3.9
		Hs.111496			
		Hs.79428	AA521047	BCL2/adenovirus E1B 19kD-interacting pro	3.9
	114555	Hs.167904	AA058594	ESTs	3.9
	122138	Hs.163960	AA435549	ESTs	3.9
35		Hs.198726		vasoactive intestinal peptide receptor 1	3.9
55		Hs.75216		protein tyrosine phosphatase; receptor t	3.9
					3.9
		Hs.325474		caldesmon 1	
	105635	Hs.301985	AA281508	ESTs	3.9
	134285	Hs.81086	AA460012	solute carrier family 22 (organic cation	3.9
40	134125	Hs.50421	R38102	KIAA0203 gene product	3.9
		Hs.241493		natural killer-tumor recognition sequenc	3.9
				ESTs	3.9
		Hs.186600			
				Caldesmon 1, Alt. Splice 6, Non-Muscle	3.9
	104334	Hs.78771	D82614	ESTs	3.9
45	110242	Hs.19978	H26417	ESTs	3.9
		Hs.289008		ESTs	3.9
		Hs.303193		zt87a9.r1 Soares_testis_NHT Homo sapiens	3.9
		Hs.293960		ESTs	3.9
		Hs.110445		ESTs; Moderately similar to unknown [M.m	3.9
50	130752	Hs.18895	D50927	KIAA0137 gene product	3.8
		Hs.112110		ESTs	3.8
		Hs.32478	AA040154	ESTs	3.8
					- 3.8
		Hs.71721	AA142913	ESTs	
		Hs.45207	AA292537	ESTs	3.8
55	100452	Hs.241552	D87742	Human mRNA for KIAA0268 gene; partial cd	3.8
	104454	Hs.129228	M84443	galactokinase 2	3.8
		Hs.102859		ESTs	3.8
				ESTs; Highly similar to (defline not ava	3.8
		Hs.24427			
		Hs.269228		ESTs	3.8
60		Hs.73848	AA069549	ESTs	3.8
	106932	Hs.9394	AA495926	ESTs	3.8
		Hs.620	M69225	bullous pemphigoid antigen 1 (230/240kD)	3.8
		Hs.14912	AA424524	Homo sapiens mRNA for KIAA0286 gene; par	3.8
~		Hs.269721		ESTs	3.8
65		Hs.58694	W92051	ESTs	3.8
	126457	Hs.50382	AA007489	zh98g04.r1 Soares_fetal_liver_spleen_1NF	3.8
		Hs.112969		EST	3.7
		Hs.17752	H95978	Homo saplens phosphatidylserine-specific	3.7
		Hs.162	M35410	insulin-like growth factor binding prote	3.7
	130377	115,102	TIOUT IV	media-liva Bioarn motor purgus bioro	U. 7

	117007	Hs.44708	N39214	ser-Thr protein kinase related to the my	3.7
				ESTs; Weakly similar to BONE/CARTILAGE P	3.7
		Hs.39712	N77278	Homo sapiens mRNA for membrane protein w	3.7
		Hs.278721			3.7
5		Hs.305971 Hs.193700		ESTs ESTs; Moderately similar to IIII ALU SUB	3.7
5				CDC16 (cell division cycle 16; S. cerevi	3.7
		Hs.1592	U18291		3.7
		Hs.183475		ESTs; Moderately similar to !!!! ALU SUB	3.7
		Hs.272531		w34b06.s1 Morton Fetal Cochlea Homo sap	3.7
10		Hs.23837 Hs.60293	N22222	ESTs	3.7
10		Hs.15683	AA496037 T92030	ESTS	3.7
		Hs.279952		ESTs; Highly similar to (defline not ava	3.7
		Hs.19347		ESTs	3.7
		Hs.291025		EST	3.7
15		Hs.22380		ESTs	3.7
13		Hs.75295	X66534	guanylate cyclase 1; soluble; alpha 3	3.7
		Hs.109390		ESTs	3.7
	_	Hs.54900		ESTs	3.7
		Hs.7337	AA512902	ESTs	3.7
20		Hs.31707		ESTs	3.7
		Hs.194283		Homo sapiens putative GR6 protein (GR6)	3.7
		Hs.35699	R97219	ESTs	3.7
		Hs.105273		ESTs	3.6
		Hs.6363	AA206625	ESTs	3.6
25				Transcription Factor lia	3.6
		Hs.166994		FAT tumor suppressor (Drosophila) homolo	3.6
	103520		Y10511	H.sapiens mRNA for CD176 protein	3.6
	113778	Hs.302738	W15263	ESTs	3.6
	101838	Hs.75511	M92934	connective tissue growth factor	3.6
30	113702		T97307	ESTs; Moderately similar to IIII ALU SUB	3.6
	118201	Hs.48428	N59800	EST	3.6
		Hs.68554	C20780	EST	3.6
•		Hs.22983	AA400517	ESTs; Moderately similar to UDP-GLUCOSE:	3.6
25		Hs.170291		ESTS	3.6 3.6
35		Hs.27973	AA806365	oc26h07.s1 NCL_CGAP_GCB1 Homo sapiens cD dioxin-responsive gene {putative polyade	3.6
	101964	Hs.326416	S81578	ESTs	3.6
		Hs.337434		ESTs	3.6
		Hs.142296		ESTs	3.6
40				V-Erba Related Ear-3 Protein	3.6
		Hs.164018		ESTs	3.6
	110758	Hs.274265	N21385	talin	3.6
	107307	Hs.44155	T52099	creatine kinase; mitochondrial 2 (sarcom	3.6
		Hs.183639		ESTs	3.6
45		Hs.184325		ESTs	3.6
		Hs.270696		ESTs; Moderately similar to !!!! ALU SUB	3.6
		Hs.199067		v-erb-b2 avian erythroblastic leukemia v	3.6
		Hs.44829		ESTs; Moderately similar to !!!! ALU SUB	3.6
50		Hs.35841	AA436459	nuclear factor I/X (CCAAT-binding transc	3.6 3.6
50		Hs.279780		ESTs	3.6
		Hs.73793	M27281	vascular endothelial growth factor	3.6
		Hs.334641		ESTs	3.5
		Hs.79572	AA235803	ESTs - arylsulfatase D	3.5
55	457951 108407		Al369384 AA075519	zm87h9.s1 Stratagene ovarian cancer (#93	3.5
33	126659		T16245	a disintegrin and metalloproteinase doma	3.5
	104189	Hs.301804		ESTs	3.5
	125056	Hs.129014	N53276	ESTs	3.5
		Hs.79386	X54162	Human mRNA for a 64 Kd autoantigen expre	3.5
60	133011	Hs.171921	AA042990	sema domain; immunoglobulin domain (lg);	3.5
	131379	Hs.26176	R49035	ESTs	3.5
	126742	Hs.169359	H64106	yr57e06.r1 Soares fetal liver spleen 1NF	3.5
	105560	Hs.306915	AA262783	ESTs	3.5
	118472	Hs.42179	N66818	ESTs	3.5
65		Hs.30127	AA280895	ESTs; Highly similar to !!!! ALU SUBFAMI	3.5
		Hs.145807		ESTs; Moderately similar to IIII ALU SUB	3.5
		Hs.26771	AA126472	ESTs	3.5
		Hs.18953		phosphodiesterase 9A	3.5 3.5
	11/4/3	Hs.155560	13013/	ESTs	لندن

		Hs.168075		karyopherin (importin) beta 2	3.5
		Hs.13531	AA442868	ESTs; Weakly similar to (defline not ava	3.5
		Hs.41119		ESTs	3.5
_	131689	Hs.30696	AA599653	transcription factor-like 5 (basic helix	3.5
5	127862	Hs.163191	AA765305	EST	3.5
	126995	Hs.189810	W26950	Human DNA sequence from PAC 388M5 on chr	3.5
	119071		R31180	ESTs .	3.5
	103941	Hs.96593	AA282978	ESTs	3.5
	110721	Hs.31319	H97678 .	ESTs	3.5
10	126586	Hs.43086	AA011247	ESTs	3.5
	103106	Hs.1857	X62025	phosphodiesterase 6G; cGMP-specific; rod	3.5
	116357	Hs.90797	AA504806	Homo sepiens clone 23620 mRNA sequence	3.5
	105309	Hs.4104	AA233790	ESTs	3.5
	130796	Hs.19525	R39390	ESTs	3.5
15	109101	Hs.52184	AA167708	ESTs	3.5
	103134	Hs.2839	X65724	Norrie disease (pseudoglioma)	3.5
	131798	Hs.301449	X86098	adenovirus 5 E1A binding protein	3.5
	118535	Hs.49418	N67968	ESTs	3.5
	102592	Hs.11223	U62389	Human putative cytosolic NADP-dependent	3.4
20	125905	Hs.6456	T69868	chaperonin containing TCP1; subunit 2 (b	3.4
	109160	Hs.301997	AA179387	ESTs	3.4
	105327	Hs.211593	AA234440	ESTs	3.4
	106586	Hs.57787	AA456598	ESTs	3.4
	122635		AA454085	EST	3.4
25	132413	Hs.260116	AA132969	metalloprotease 1 (pitrilysin family)	3.4
	131938	Hs.34956	AA283620	ESTs	3.4
	133871	Hs.182793	AA454597	ESTs	3.4
	107175	Hs.292503	AA621751	ESTs; Weakly similar to KIAA0601 protein	3.4
	101188	Hs.184298	L20320	cyclin-dependent kinase 7 (homolog of Xe	3.4
30	126422	Hs.237658	H48518	ESTs; Highly similar to apolipoprotein A	3.4
	118475		N66845	ESTs; Weakly similar to IIII ALU CLASS B	3.4
	104558	Hs.88959	R56678	ESTs; Weakly similar to !!!! ALU SUBFAMI	3.4
	128307	Hs.132005	Al453794	ESTs	3.4
	112254	Hs.25829	R51831	ESTs	3.4
35	125408	Hs.89578	N72353	yv37e12.r1 Soares fetal liver spleen 1NF	3.4
	109834	Hs.175955	H00604	ESTs	3.4
	130844	Hs.20191	D12122	seven in absentia (Drosophila) homolog 2	3.4
	127143	Hs.20843	AA533553	nj68h04.s1 NCI_CGAP_Pr10 Homo saplens cD	3.4
	135309	Hs.42500	D25984	ESTs	3.4
40	125724	Hs.295978	AA083407	stimulated trans-acting factor (50 kDa)	3.4
	127692	Hs.187983	Al021912	ESTs	3.4
	116674	Hs.92127	F04816	ESTs	3.4
	134700	Hs.8868	AA481414	golgi SNAP receptor complex member 1	3.4
		Hs.166196		ESTs	3.4
45		Hs.155983		H.sapiens mRNA for 5'UTR for unknown pro	3.4
		Hs.89925		calcium channel; voltage-dependent; L ty	3.4
		Hs.16085	AA232535	ESTs; Highly similar to (defline not ava	3.4
	111331	Hs.15978	N78773	ESTs	3.4
~~		Hs.10653	AA412505	ESTs	3.4
50		Hs.21893	R45698	ESTs	3.4
		Hs.35828		ESTs	3.4
		Hs.255015		ob93c10.s1 NCI_CGAP_GCB1 Homo sapiens cD	3.4
		Hs.75616		KIAA0018 gene product	3.4
		Hs.247992		Homo sapiens DNA binding protein for sur	3.4
55		Hs.20621	T08287	ESTs	3.4
		Hs.26994	AA489009	ESTs	3.4
		Hs.302267		ESTs	3.4
	125957		H45213	yo03b08.r1 Soares adult brain N2b5HB55Y	3.3
CO		Hs.146085		ESTs	3.3
60		Hs.75354	F13702	ESTs	3.3
		Hs.170098		ESTs; Highly similar to KIAA0372 [H.sapi	3.3
		Hs.143792	_	ESTs; Weakly similar to glioma amplified	3.3
		Hs.75454	D49396	Human mRNA for Apo1_Human (MER5(Aop1-Mou	3.3
15		Hs.30692	U24153	p21 (CDKN1A)-activated kinase 2	3.3
65		Hs.88201	AA481256	ESTs; Weakly similar to (defline not ava	3.3
	102034		U05291	fibromodulin	3.3
		Hs.14658	R99606	Human chromosome 5q13.1 clone 5G8 mRNA	3.3
		Hs.159456		ESTs; Highly similar to (defline not ava	3.3
	126707	Hs.104105	MA1304/4	Meis (mouse) homolog 2	3.3

	445010	II- 400057	4.4020000	COT-	
		Hs.190057		ESTS	3.3
		Hs.31110		ESTs	3.3
		Hs.24192	R31679	ESTs	3.3
_		Hs.2877	X63629	cadherin 3; P-cadherin (placental)	3.3
5		Hs.62604		ESTs	3.3
		Hs.79284		mesoderm specific transcript (mouse) hom	3.3
		Hs.301804		ESTs	3.3
	102165	Hs.159627	U18321	Death associated protein 3	3.3
	126966	Hs.182575	R38438	solute carrier family 15 (H+/peptide tra	3.3
10	124839	Hs.140942	R55784	ESTs	3.3
	100709	Hs,100469	HG3264-HT3441	Af-6 (Gb:U02478)	3.3
	132967	Hs.61635	AA032221	Homo sapiens BAC clone RG041D11 from 7q2	3.3
		Hs.65114		keratin 18	3.3
	132616	Hs.283558		ESTs	3.3
15		Hs.129781		ESTs	3.3
		Hs.31652		ESTs	3.3
		Hs.87113		ESTs	3.3
		Hs.112227		ESTs	3.3
		Hs.12315		ESTs	3.3
20		Hs.178604		ESTs	3.3
20		Hs.155995		Homo sapiens mRNA for KIAA0643 protein;	3.3
		Hs.284294		ESTs	3.3
		Hs.7569	T26893	EST	3.3
		Hs.82318		Brush-1	3.3
25		Hs.333256		ESTs; Moderately similar to !!!! ALU SUB	3.3
25				ESTs	3.3
		Hs.294105			3.3
		Hs.194215		ESTS	3.3
		Hs.299867		hepatocyte nuclear factor 3; alpha	
20		Hs.190151		ESTS	3.3
30		Hs.47402		ESTs; Weakly similar to !!!! ALU SUBFAMI	3.3
		Hs.11500	AA437118	ESTS	3.3
		Hs.126494		ESTS	3.3
	127265		AA332751	EST37214 Embryo, 8 week I Homo sapiens c	3.3
25		Hs.41143		Homo sapiens mRNA for KIAA0581 protein;	3.2
35		Hs.293691		ESTs	3.2
		Hs.250655		H.sapiens mRNA for Ptg-12 protein	3.2
		Hs.334334		ESTs	3.2
		Hs.251946		ESTs	3.2
40		Hs.44481		forkhead (Drosophila)-like 6	3.2
40		Hs.32425	AA488889	ESTs	3.2
		Hs.169780		homologous to yeast nitrogen permease (c	3.2
		Hs.292581		ESTs	3.2
		Hs.284207		ESTs	3.2
		Hs.105116		EST	3.2
45		Hs.63908	AA598745	ESTs	3.2
		Hs.194657		H.sapiens gene encoding E-cadherin, exon	3.2
	124006	Hs.270016	D60302	ESTs	3.2
		Hs.74137	L40391	Homo sapiens (clone s153) mRNA fragment	3.2
	101183	Hs.795	L19779	H2A histone family; member O	3.2
50	125596		R25698	yg44h11.r2 Soares infant brain 1NIB Homo	3.2
	127261		AA661567	nu86b02.s1 NCI_CGAP_Aiv1 Homo sapiens cD	3.2
	120090	Hs.59554	W94591	ESTs	3.2
	129393	Hs.166982	D13435	phosphatidylinositol glycan; class F -	3.2
	120923	Hs.97129	AA382283	ESTs	3.2
55	118907	Hs.274256	N91003	ESTs	3.2
	111552	Hs.191185	R09411	ESTs	3.2
	104431	Hs.99913	J03019	adrenergic; beta-1-; receptor	3.2
		Hs.278634	D63480	Human mRNA for KIAA0146 gene; partial cd	3.2
	131615	Hs.192803	D14533	xeroderma pigmentosum; complementation g	3.2
60		Hs.84072	U47732	transmembrane 4 superfamily member 3	3.2
		Hs.116774		Integrin; alpha 1	3.2
		Hs.24095	W68845	ESTs	3.2
		Hs.70937	Z83735	H3 histone family; member K	3.2
		Hs.189716		ESTs	3.2
65		Hs.104696		ESTs .	3.2
33		Hs.6639	W28406	ESTs	3.2
		Hs.334335		ESTs	3.2
		Hs.185766		ESTs	3.2
		Hs.130760		Homo sapiens mRNA; cDNA DKFZp586N0318 (f	3.2
	101012		47000110	Linuin adhigiis ilii ilitri emitri piii efeccitorio (i	J.E.

	102214	Hs.32964	U23752	SRY (sex-determining region Y)-box 11	3.2
	123147		AA487961	ab11h6.s1 Stratagene lung (#93721) Homo	3.2
		Hs.272138	R00940	ye87g03.r1 Soares fetal liver spleen 1NF	3.2
	116246	Hs.250646	AA479961	ESTs; Highly similar to ubiquitin-conjug	3.2
5		Hs.180789		Homo sapiens (clone S164) mRNA; 3' end o	3.2
_			AF001548	myosin; heavy polypeptide 11; smooth mus	3.2
		Hs.304389		ESTs	3.2
		Hs.67619	AA089559	Homo sapiens mRNA; chromosome 1 specific	3.2
		Hs.182378		colony stimulating factor 1 (macrophage)	3.2
10		Hs.242894		ADP-ribosylation factor-like 1	3.1
		Hs.234896		ESTs; Highly similar to geminin [H.sapie	3.1
		Hs.5669	C14290	ESTs	3.1
	123926	Hs.227933	AA621348	ESTs; Highly similar to (defline not ava	3.1
		Hs.239720		ESTs; Weakly similar to Rga [D.melanogas	3.1
15	105322	Hs.16346	AA234100	ESTs	3.1
	100631	Hs.48332	HG2709-HT2805	Serine/Threonine Kinase (Gb:Z25431)	3.1
	130791	Hs.199263	AA259102	ESTs; Highly similar to (defline not ava	3.1
	131220	Hs.300855	R77200	ESTs	3.1
	113237	Hs.123642	T62857	ESTs	3.1
20	125562	Hs.98968	Al494372	ESTs	3.1
	134110	Hs.79136	U41060	Human breast cancer; estrogen regulated	3.1
		Hs.47334	W85888	ESTs; Moderately similar to !!!! ALU SUB	3.1
		Hs.296842		ESTs; Moderately similar to non-muscle m	3.1
~~		Hs.40719	AA299096	Homo sapiens mRNA; cDNA DKFZp564M0916 (f	3.1
25		Hs.286192		ESTs	3.1
		Hs.296141		ESTs .	3.1
		Hs.178294		ESTs	3.1
		Hs.183297		ESTS	3.1
30		Hs.109653		ESTs	3.1 3.1
30		Hs.90800		matrix metalloproteinase 16 (membrane-in	3.1
		Hs.38972 Hs.182828	AA161043	tetraspan 1 zinc finger protein 136 (clone pHZ-20)	3.1
		Hs.33010	H80622	Homo sapiens mRNA for KIAA0633 protein;	3.1
		Hs.44698	N35115	ESTs	3.1
35		Hs.287849		ESTs	3.1
-		Hs.103120		ESTs	3.1
	100789			Phosphoglucomutase 1, Alt. Splice	3.1
	126017	Hs.159440	H60487	ESTs	3.1
	132452	Hs.247324	AA005262	Homo sapiens DNA sequence from PAC 262D1	3.1
40		Hs.108479		ESTs	3.1
		Hs.181368		U5 snRNP-specific protein (220 kD); orth	3.1
		Hs.118258		ESTs	3.1
	123465		AA599033	ESTS	3.1
15		Hs.152316		EST51345 Gall bladder II Homo sapiens cD	3.1
45		Hs.167031		za36d05.r1 Soares fetal liver spleen 1NF	3.1 3.1
		Hs.43234	N72094	ESTs	3.1
		Hs.38057	AA203742	ESTs	3.1
		Hs.124347 Hs.223241		ESTs yb15c11.s1 Stratagene placenta (#937225)	3.1
50		Hs.15220	AA249334	i312.seq.F Human fetal heart, Lambda ZAP	3.1
50		Hs.22242	AA463737	ESTs.	3.1
		Hs.20993		ESTs; Weakly similar to Ydr374cp [S.cere	3
		Hs.35014	W92779	ESTs -	3
		Hs.106390		ESTs	3
55		Hs.247815		H.sapiens H4/I gene	3
		Hs.250614		yy13h06.r1 Soares melanocyte 2NbHM Homo	3
		Hs.21275	D25755	ESTs	3
		Hs.75354		ESTs	3
		Hs.5811	R12421	ESTs	3
60	127363	Hs.22116	AA307744	Homo sapiens Cdc14B1 phosphatase mRNA; c	3
	103690	Hs.84063	AA016186	ESTs	3
	102589	Hs.8867	U62015	Homo sapiens Cyr61 mRNA, complete cds	3
	125144	Hs.24336	W37999	ESTs	3
	132977	Hs.301404	U28686	RNA binding motif protein 3	3
65		Hs.146170		ESTs	3
		Hs.79411	J05249	replication protein A2 (32kD)	3
		Hs.248177		Human histone H3 gene	3
		Hs.30738	AA257971	ESTs	3
	13113/	Hs.33287	U85193	nuclear factor I/B	3

	407004	U= 044554	A1054000	FOT-	•
		Hs.241551		ESTS	3 3
		Hs.24104 Hs.3066	R26708 U26174	ESTs granzyme K (serine protease; granzyme 3;	3
		Hs.21291		Serine/Threonine Kinase (Gb:Z25428)	3
5			W86838	EST COMMO (CIMASO (CIMASO)	3
5		Hs.118281		zinc finger protein 266	3
		Hs.76152	M14219	decorin	3
		Hs.14449	AA010889	ESTs	3
		Hs.304139		EST	3
10	127635	Hs.116346	AA766903	ESTs	3
	128434	Hs.143880	Al190914	ESTs	3
		Hs.187555		ESTs	- 3
				ESTs	3 3 3
٠,٠		Hs.103804		heterogeneous nuclear ribonucleoprotein	. 3
15		Hs.251531		proteasome (prosome; macropain) subunit;	3
			AA621125	Homo sapiens chromosome 2; 10 repeat reg	3
		Hs.22545 Hs.263727	R43910	ESTs ESTs; Moderately similar to !!!! ALU SUB	3
•		Hs.21739	AA370302	Homo sapiens mRNA; cDNA DKFZp58611518 (f	3
20		Hs.274407		thymus specific serine peptidase	3
20		Hs.75847	N79435	ESTs	3
		Hs.227949		ESTs; Highly similar to SEC13-RELATED PR	3
		Hs.44189	N30426	ESTs	3
	123659	Hs.112699	AA609368	ESTs	3
25 ·	103963	Hs.63290	AA298588	EST114219 HSC172 cells II Homo sapiens c	3
		Hs.7367	AA112222	ESTs; Moderately similar to (defline not	3
•		Hs.80975	AA255903	CD39-like 4	2.9
		Hs.89890	S72370	pyruvate carboxylase	2.9
20		Hs.101810		ESTs; Weakly similar to !!!! ALU SUBFAMI	2.9 2.9
30		Hs.7980 Hs.112575	F09570	ESTs .	2.9
		Hs.32793	AA609943	ESTs	2.9
		Hs.88556	D50405	histone deacetylase 1	2.9
		Hs.82689	H99801	tumor rejection antigen (gp96) 1	2.9
35		Hs.174139		H. sapiens RNA for CLCN3	2.9
		Hs.14512	AA461495	ESTs	2.9
	108555		AA084963	zn13e12.s1 Stratagene hNT neuron (#93723	2.9
		Hs.2110	HG945-HT945	Nucleic Acid-Binding Protein (Gb:L12693)	2.9
40		Hs.16492	AA173998	ESTs; Weakly similar to weakly similar t	2.9
40		Hs.139226		replication factor C (activator 1) 2 (40	2.9
	106636		AA459950	ESTs	2.9 2.9
		Hs.108708 Hs.251871		calcium/calmodulin-dependent protein kin stromal cell-derived factor 1	2.9
		Hs.9857	AA433946	ESTs; Weakly similar to (defline not ava	2.9
45		Hs.301636		peroxisomal biogenesis factor 6	2.9
		Hs.98074	AA056263	ESTs; Moderately similar to IIII ALU SUB	2.9
		Hs.9701	AA402224	Homo sapiens growth arrest and DNA-damag	2.9
	108552		AA084912	zn11c7.s1 Stratagene hNT neuron (#937233	2.9
	126505	Hs.190057	W26894	16a11 Human retina cDNA randomly primed	2.9
50		Hs.79086	X06323	Human MRL3 mRNA for ribosomal protein L3	2.9
		Hs.211539		eukaryotic translation initiation factor	2.9
		Hs.277422		Homo sapiens mRNA for cadherin FIB3, par	2.9
		Hs.44104		ESTs ESTs; Moderately similar to WAP four-dis	- 2.9 2.9
55		Hs.36688 Hs.250870		protein kinase; mitogen-activated; kinas	2.9
<i>JJ</i>		Hs.32995		ESTs	2.9
		Hs.26267		ESTs; Weakly similar to torsinA [H.saple	2.9
		Hs.173824		thymine-DNA glycosylase	2.9
	100641	Hs.182183	HG2743-HT2846	Caldesmon 1, Alt. Splice 4, Non-Muscle	2.9
60	104602		R86920	ESTs	2.9
•		Hs.42738		ESTs	2.9
		Hs.34073		BH-protocadherin (brain-heart)	2.9
•		Hs.155212		methylmalonyl Coenzyme A mutase	2.9
65		Hs.5724	AA279422	ESTs	2.9
65		Hs.287912		lectin; mannose-binding; 1	2.9 2.9
		Hs.19762		ESTs H.sapiens mRNA for ArgBPIB protein	2.9
		Hs.285728 Hs.132390	_	ESTs	2.9
•		Hs.161002		absent in melanoma 1	2.9
		. 1911 9 1946			•

	131710	Hs.30985	AA233225	ESTs; Highly similar to (defline not ava	2.9
		Hs.268903		ESTs	2.9
		Hs.15535	Al417137	Homo sapiens clone 24582 mRNA sequence	2.9
	104229	Hs.61289	AB002346	inositol phosphate 5'-phosphatase 2 (syn	2.9
5	126600	Hs.191385	AA699949	ESTs	2.9
	125175	Hs.303030	W52355	EST	2.9
	103849	Hs.34578	AA187045	ESTs; Weakly similar to IIII ALU SUBFAMI	2.9
		Hs.78961	U14575	protein phosphatase 1; regulatory (inhib	2.9
10		Hs.107815		ESTs	2.9
10		Hs.303125		ESTs	2.9
		Hs.218329		heat shock 70kD protein 1	2.9
		Hs.75462	U72649	Human BTG2 (BTG2) mRNA; complete cds	2.9 2.9
		Hs.18271	AA191014	ESTs; Wealdy similar to Ydr372cp [S.cere Human mRNA for transcription factor AREB	2.9
15		Hs.232068 Hs.336901		ESTs	2.9
13		Hs.37637	N59645	ESTS	2.9
		Hs.11805	N66066	ESTs	2.9
		Hs.102897		ESTs	2.9
		Hs.79265	AA114183	ESTs; Moderately similar to glutamate py	2.9
20		Hs.267812		sorting nexin 4	2.9
		Hs.279609		pigment epithelium-derived factor	2.9
	103803	Hs.106149	AA127696	ESTs	2.9
	102139	Hs.2128	U15932	dual specificity phosphatase 5	2.9
	128104		AA971000	op67g11.s1 Soares_NFL_T_GBC_S1 Homo sapi	2.8
25		Hs.337631		nz22d08.s1 NCI_CGAP_GCB1 Homo sapiens cD	2.8
		Hs.180952		ESTs	2.8
		Hs.217916		ESTs (Observable of Total)	2.8
		Hs.93883	D10537	myelin protein zero (Charcot-Marie-Tooth	2.8 2.8
20		Hs.68644	N45014	yy80g06.r1 Soares_multiple_sclerosis_2Nb	2.8
30		Hs.145696 Hs.98684		ESTs ESTs	2.8
		Hs.322645		ESTs	2.8
		Hs.99915		androgen receptor (dihydrotestosterone r	2.8
		Hs.129998		ESTs	2.8
35		Hs.109019		ESTs	2.8
-		Hs.12186	R45480	cyclin K	2.8
		Hs.32149	H45968	ESTs	2.8
		Hs.5409	AF008442	RNA polymerase I subunit	2.8
	105366	Hs.282093	AA236356	ESTs	2.8
40		Hs.5957	AA417761	Homo sapiens clone 24416 mRNA sequence	2.8
		Hs.25960	M13241	v-myc avian myelocytomatosis viral relat	2.8
		Hs.26255	R42714	EST	2.8
		Hs.250175		Homo sapiens clone 23904 mRNA sequence	2.8 2.8
45		Hs.33130	H44825 AA236843	ESTs ESTs; Weakly similar to unknown [S.cerev	2.8
43		Hs.72085	R20353	vg20f10.r1 Soares infant brain 1NIB Homo	2.8
	128152	Hs.23740		ESTs	2.8
		Hs.97101	AA215333	ESTs	2.8
		Hs.184510		stratifin	2.8
50		Hs.293845		ESTs	2.8
••		Hs.292566		ESTs	2.8
		Hs.12372	H98141	ESTs	2.8
		Hs.98541	AA282787	ESTs; Highly similar to (defline not ava	2.8
		Hs.46847	AA256524	Human DNA sequence from clone 30M3 on ch	2.8
55	133865	Hs.170290	F09315	discs; large (Drosophila) homolog 5	2.8
		Hs.110826		Homo sapiens CAGF9 mRNA; partial cds	2.8
		Hs.23767	Z38910	ESTs	2.8
		Hs.10299	H09594	ESTs; Moderately similar to IIII ALU SUB	2.8
6 0		Hs.66731	U81599	homeo box B13	2.8
60		Hs.336629 Hs.25067	R49116	ESTs; Weakly similar to zinc finger prot EST	2.8 2.8
	112100	Hs.20007			2.8
		Hs.173694 Hs.6019	AA430108	ESTs; Moderately similar to (defline not ESTs	2.8
	121060	Hs.22564	AA160890	myosin VI	2.8
65	111383	Hs.40919	N94527	ESTs	2.8
0 5		Hs.1594	U14518	centromere protein A (17kD)	2.8
		Hs.79981	U79242	Human clone 23560 mRNA sequence	2.8
	129887	Hs.274324	W92041	PCAF associated factor 65 alpha	2.8
		Hs.181297		ESTs	2.8

	104367	Hs.134342	H17438	ESTs; Weakly similar to seventransmembra	2.8
		Hs.193700		ESTs; Moderately similar to IIII ALU SUB	2.8
	128059	Hs.145096	AA972446	ESTs	2.8
	124447		N48000	ESTs	2.8
_					
5	111398	Hs.125565	H00086	deafness; X-linked 1; progressive	2.8
	134085	Hs.79018	U20979	chromatin assembly factor I (150 kDa)	2.8
					2.8
	124788	Hs.100912	H43543	ESTs	
	112248	Hs.326416	R51361	ESTs	2.8
					2.8
	121309	Hs.97312	AA402482	ESTs	
10	103076	Hs.75319	X59618	ribonucleotide reductase M2 polypeptide	2.8
					2.8
	10/0/1	Hs.35198	AA609053	ESTs	
	104425	Hs.35380	H88496	ESTs ·	2.8
		Hs.62245	AA446906	solute carrier family 25 (mitochondrial	2.8
	104968	Hs.29669	AA084602	ESTs	2.8
15	121153	Hs.97694	AA399640	ESTs	2.8
10					
	131216	Hs.243901	D31008	ESTs	2.8
	109682	Hs.22869	F09299	ESTs	2.8
				ESTs; Moderately similar to roundabout 1	2.8
		Hs.168818			
	132027	Hs.181444	N78844	ESTs; Weakly similar to R12C12.6 [C.eleg	2.8
20		Hs.190478		ESTs	2.8
20					
	132598	Hs.530	M81379	collagen; type IV; alpha 3 (Goodpasture	2.8
	101121	Hs.1313	L09753	tumor necrosis factor (ligand) superfami	2.8
	123000	Hs.105640	AA479347	ESTs	2.8
		Hs.1755	AA404324	ESTs ·	2.8
25					2.7
25	100481	HS.121489	HG1098-HT1098	Cystatin D	-
	113803	Hs.283683	W42789	ESTs	2.7
				ESTs; Weakly similar to cytochrome P-450	2.7
	110934	Hs.169001			
	432888		T86823	ESTs	2.7
		Hs.188898	ANADARON	ESTs	2.7
20					
30	130396	Hs.155313	AB002331	Human mRNA for KIAA0333 gene; partial cd	2.7
	121103	Hs.97697	AA398936	ESTs; Weakly similar to (defline not ava	2.7
					2.7
	131129	Hs.23240	R27296	ESTs	
	130943	Hs.272429	D50855	calcium-sensing receptor (hypocalciuric	2.7
		Hs.87819		ESTs; Weakly similar to keratin 9; cytos	2.7
0.5					
35	111900	Hs.25318	R39044	ESTs	2.7
_	106025	Hs.173334	AA412063	ESTs	2.7
	126144	Hs.40639	N39696	yx92a07.r1 Soares melanocyte 2NbHM Homo	2.7
	103248	Hs.75262	X77383	cathepsin O	2.7
					2.7
	12/230	Hs.274170	LISUSUI	Homo saplens Opa-Interacting protein OIP	
40	101584	Hs.84072	M35252	transmembrane 4 superfamily member 3	2.7
. •				ESTs	2.7
		Hs.167489			
	129689	Hs.77873	AA130156	ESTs	2.7
		Hs.9973	W92797	ESTs	2.7
	120827	Hs.132967	AA347717	ESTs ·	2.7
45	12/570	Hs.85963	N23222	ESTs; Moderately similar to IIII ALU SUB	2.7
1 5					
		Hs.256301		ESTs	2.7
	132037	Hs.332541	AA203649	ESTs; Weakly similar to HEM45 [H.sapiens	2.7
					2.7
		Hs.179825		Human sperm membrane protein BS-63 mRNA,	
	122851	Hs.99598	AA463627	ESTs	2.7
50		Hs.196384		prostaglandin-endoperoxide synthase 2 (p	2.7
50					
	120537	Hs.160422	AA262790	ESTs	2.7
		Hs.174140		ATP citrate lyase	2.7
	133889	Hs.211582	Teseevaa	ESTs	- 2.7
		Hs.106529		zv81e01.r1 Soares_total_fetus_Nb2HF8_9w	2.7
55					2.7
55		Hs.306044		ESTs	
	423239		AA323591	EST26392 Cerebellum II Homo sapiens cDNA	2.7
		Hs.12321		ESTs	2.7
	126021	Hs.187516	AA775894	ESTs	2.7
	102116		U13706	Human ELAV-like neuronal protein 1 isofo	2.7
60		11- 007007			
60	133394	Hs.237225	W10/28	ESTs; Weakly similar to (defline not ava	2.7
	104267	Hs.278439	C00358	ESTs	2.7
					2.7
		Hs.40241		ESTs; Highly similar to (defline not ava	
	129809	Hs.1259	X55283	asialoglycoprotein receptor 2	2.7
				ESTs: Weakly similar to !!!! ALU SUBFAMI	2.7
~~		Hs.283309			
65	128422		T85681	yd60c06.r1 Soares fetal liver spleen 1NF	2.7
		Hs.43899		ESTs	2.7
	118696	Hs.292284	N/2086	Homo sapiens RNA polymerase III largest	2.7
	106053	Hs.36727	AA416963	ESTs; Highly similar to histone H2A [H.s	2.7
					2.7
	104440	Hs.284380	1.20492	gamma-glutamyltransferase 1	2.1

129426 Hs.111323 AA412087 EST; Highly similar to (defline not ava					The state of the s	
106716 hs.238928 A.464962 ESTS 106863 278291 578291			Hs.111323		EST; Highly similar to (defline not avai	2.7
103963						_
114162 Hs.22665 738498 ESTS 173498 ESTS 173897 AA773857 173498 ESTS 173897 AA773857 173897 AA773857 173897 AA773857 173897 AA773857 173896 Hs.16803 AA621718 ESTS; Highly similar to (defline not ava 2.7 AA773857 173896 Hs.246358 A26388 A26388 A26388 A26388 A26388 AA79388 ESTS; Highly similar to (defline not ava 2.7 A26388 A263		106716	Hs.238928	AA464962		
119063 H.S.027 T32438 ESTS 217897 AA773857 AA773857 AA773857 ESTS; Weakly similar to (defline not ava 2.7 AA78958 ESTS; Weakly similar to (defline not ava 2.7 AA78958 ESTS; Weakly similar to (defline not ava 2.7 AA78958 ESTS; Weakly similar to (defline not ava 2.7 AA79958 ESTS; Weakly similar to (defline not ava 2.7 AA79958 ESTS; Weakly similar to (defline not ava 2.7 AA79958 ESTS; Weakly similar to Similar to colla 2.7 AA79958 ESTS; Weakly similar to Similar to colla 2.7 AA79958 ESTS; Weakly similar to Similar to colla 2.7 AA79958 ESTS; Weakly similar to Similar to colla 2.7 AA79958 ESTS; Weakly similar to (defline not ava 2.7 AA79958 ESTS; Moderately similar to MAA0350 AA79958 ESTS; Meakly similar to MAA0350 AA79958		103663		Z78291	Z78291 Homo saplens brain fetus Homo sap	2.7
197697	5	114162	Hs.22265	Z38909	ESTs	2.7
130821 Hs.18803 AA621718 ESTs; Weakly similar to (define not ava 2.7		113063	Hs.5027	T32438	ESTs	2.7
16245 Hs.42706 AA79988 ESTS; Highly similar to (defline not ava 2.7		127897		AA773857	af80c09.r1 Soares_NhHMPu_S1 Homo sapiens	2.7
116245 Is.42786 AA79988 ESTs; Highly similar to (delline not ava 2.7 133800 Is.77899 M19297 tropomyosin 1 (alpha) 2.7 134902 Is.92303 A268043 ESTs; Weakly similar to Similar to colla 2.7 15 106803 Is.284285 AA4054726 ESTs 2.7 16 106803 Is.284285 AA4054726 ESTs 2.7 17 1253 Is.15788 AX054726 ESTs 2.7 18449 Is.164478 N66413 ESTs; Moderately similar to (IIIAU) SUB 2.7 18449 Is.164478 N66413 ESTs; Moderately similar to (IIIAU) SUB 2.7 18459 Is.984318 M63488 ESTs; Weakly similar to (IIIAU) SUB 2.7 18459 Is.984318 M63488 ESTs; Weakly similar to (IIIAU) SUB 2.7 183805 Is.24775 W56590 ESTs; Weakly similar to (IIIAU) SUB 2.7 183902 Is.24775 W56590 ESTs; Weakly similar to (IIIAU) SUB 2.7 183902 Is.24775 W56590 ESTs ESTs; Weakly similar to (IIIAU) SUB 2.7 183902 Is.24775 W56590 ESTs ESTs; Weakly similar to (IIIAU) SUB 2.7 183903 Is.26864 AA468072 ESTs ESTs; Weakly similar to (IIIAU) SUB 2.7 183904 Is.2686 AA488072 ESTs ESTs 2.7 183904 Is.2687 AA488072 ESTs ESTs 2.7 183904 Is.2687 AA488072 ESTs EST		130621	Hs.16803	AA621718	ESTs; Weakly similar to (defline not ava	2.7
125499						2.7
133860 Hs.77889 M19267 Iropomyosin 1 (alpha) 2.7 134982 Hs.92308 N48086 ESTS 2.7 2.5	10					2.7
104470 Hs.246358 N.28843 ESTs; Weakly similar to Similar to colla 2.7			Hs.77899			2.7
134882 hs.92308 N.46086 ESTs 2.7						2.7
106803 Hs.284295 AAA79114						2.7
104899						2.7
12540 Hs.337585 AB204637	15					
111253 Hs. 15768 N70042 ESTs; Moderately similar to (III) ALU SUB 2.7						2.7
118449 Hs.164478 M66413 ESTs; Weakly simillar to (defiline not ava 2.7						
134507 Hs.94318 M63498 replication protein A1 (70kD) 2.7					• • • • • • • • • • • • • • • • • • •	
121609						
118835 Hs.27475 W56590 ESTs; Highly similar to (defline not ava	20					
13962 Hs.285290 W86375	20					
121913 Hs.98558 AA428062 ESTs 2.7						
25						
2.7 130799 Hs.12695 AA484273 ESTs L23184 Hs.18166 AA489072 Homo saplens mRNA for KIAA0870 protein; 2.7 106186 Hs.6315 AA427398 Acetylserotonin N-methyltransferase-like 2.7 106186 Hs.6315 T16559 ESTs L26181 Hs.26640 Al283162 L33054 Hs.291079 R07876 ESTs; Weakly similar to unknown [S.cerev 2.7 L3131 Hs.25640 Al283162 L32630 R40752 ESTs Weakly similar to unknown [S.cerev 2.7 L3131 Hs.25640 Al283162 L32630 R40752 ESTs Weakly similar to unknown [S.cerev 2.7 L32131 Hs.25640 Al283162 ESTs Weakly similar to unknown [S.cerev 2.7 L36507 Hs.23964 Al382218 ESTs Weakly similar to unknown [S.cerev 2.6 L36507 Hs.23964 Al382218 ESTs Weakly similar to unknown [S.cerev 2.6 L36507 Hs.23964 Al382218 ESTs Weakly similar to unknown [S.cerev 2.7 L36507 Hs.23964 Al382218 ESTs Weakly similar to unknown [S.cerev 2.7 L36507 Hs.47111 Ns0740 ESTs Weakly similar to unknown [S.cerev 2.7 L36507 Hs.47111 Ns0740 ESTs Weakly similar to unknown [S.cerev 2.7 L36507 Hs.47111 Ns0740 ESTs Weakly similar to unknown [S.cerev 2.7 L36507 Hs.47111 Ns0740 ESTs Weakly similar to unknown [S.cerev 2.7 L36507 L36507 Hs.47111 Ns0740 ESTs Weakly similar to unknown [S.cerev 2.7 L36507 L36507 Hs.47111 Ns0740 ESTs Weakly similar to unknown [S.cerev 2.7 L36507 L36507 Hs.47111 Ns0740 ESTs William						
123184 Hs.18166 AA489072 Homo saplens mRNA for KIAA0870 protein; 2.7 103420 Hs.173497 X97065 SEC23-like protein B 2.7 10349 L77559 Homo saplens DGS-B partial mRNA 2.7 113945 Hs.6655 T16559 ESTS ESTS; Weakly similar to unknown [S.cerev 2.7 128131 Hs.25640 AL283162 Claudin 3 2.6 101864 Hs.75777 M85787 ESTS; Weakly similar to unknown [S.cerev 2.7 111948 Hs.26303 R40752 ESTS 2.6 117930 Hs.47111 N50740 ESTS 2.6 117930 Hs.47111 N50740 ESTS 2.6 117930 Hs.47111 N50740 ESTS 2.6 132227 Hs.4248 AA412620 ESTS 2.6 132574 Hs.274256 H03574 H036754 ESTS 2.6 133574 Hs.19303 AA456887 ESTS 2.6 1330792 Hs.19500 AA307896 Homo saplens unknown protein mRNA, parti 2.6 1330792 Hs.19500 AA307896 Homo saplens unknown protein mRNA, parti 17477 Hs.4475 M30328 ESTS 2.6 130792 Hs.19500 AA307896 nuclear localization signal deleted in v 2.6 13367 Hs.194364 AA435542 ESTS 2.6 130792 Hs.19500 AA307896 nuclear localization signal deleted in v 2.6 130792 Hs.19500 AA307896 ESTS 2.6 130792 Hs.19503 AA456687 ESTS 2.6 130792 Hs.19503 AA456687 ESTS 2.6 130792 Hs.19523 CO2582 ESTS 2.6 130792 Hs.19623 CO2582 ESTS 2.6 130792 Hs.19623 CO2582 ESTS 2.6 130792 Hs.19623 CO2582 ESTS 2.6 130793 Hs.19623 CO2582 ESTS 2.6 130793 Hs.19623 AA478429 ESTS 2.6 130794 Hs.19623 AA478429 ESTS 2.6 130794 Hs.19623 AA478429 ESTS 2.6 130968 Hs.19430 AA478429 ESTS 2.6 130968 Hs.19430 AA478429 ESTS ESTS 2.6 130968 Hs.19430 AA478429 E	25					
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121/10 MS.96/44 AA419011 ESTS 2.6						
		121710	MS.96744	AA418U11	E315	2.6

	125428	Hs.851	W74608	ESTs; Highly similar to (defline not ava	2.6
	115906	Hs.82302	AA436616	ESTs	2.6
	108432		AA076626	Homo sapiens clone 23851 mRNA sequence	2.6
	126191	Hs.191911	H97728	ESTs	2.6
5		Hs.281434		ESTs	2.6
,		Hs.268615		ESTs	2.6
		Hs.173840		ESTs	2.6
		ns.173040	U59748		2.6
	102565	11 40400	•••	Human desert hedgehog (hDHH) mRNA, parti	
10		Hs.13109	AA194973	ESTs	2.6
10		Hs.334609		ESTs	2.6
		Hs.21104		ESTs	2.6
	135192	Hs.321709	AF000234	purinergic receptor P2X; ligand-gated io	2.6
	109833	Hs.29889	H00580	ESTs	2.6
	105756	Hs.8535	AA303088	ESTs; Weakly similar to transformation-r	2.6
15		Hs.97967	AA406210	ESTs	2.6
		Hs.155485		Human huntingtin interacting protein (HI	2.6
		Hs.102329		ESTs	2.6
		Hs.97199		ESTs	2.6
	127021	Hs.180591	D88383	ESTs; Weakly similar to weak similarity .	2.6
20				ESTs; Weakly similar to (define not ava	2.6
20		Hs.11463			2.6
		Hs.26904	R61680	ESTs	2.6
		Hs.112981		ESTs	
		Hs.104835		ESTs	2.6
~ ~		Hs.110637		Homo sapiens homeobox protein A10 (HOXA1	2.6
25		Hs.153934		core-binding factor; runt domain; alpha	2.6
	130021	Hs.1435	M24470	guanosine monophosphate reductase	2.6
	100585	Hs.199160	HG2367-HT2463	Trithorax Homolog Hrx	2.6
	104965	Hs.30177	AA084104	ESTs	2.6
		Hs.46485		EST	2.6
30	124792	Hs.48712	R44357	ESTs	2.6
		Hs.74313		ESTs	2.6
		Hs.32971		phosphoinositide-3-kinase; class 3	2.6
		Hs.195614		KIAA0017 gene product	2.6
		Hs.169977		ESTs	2.6
35		115.105577		Forkhead Family Afx1	2.6
33	100858	H- 001007			2.6
		Hs.301927		T-cell receptor; alpha (V;D;J;C)	
		Hs.133865		ESTs	2.6
		Hs.92137		v-myc avian myelocytomatosis viral oncog	2.6
40		Hs.10247		Human (memc) mRNA, 3'UTR	2.6
40		Hs.191538		ESTs	2.6
		Hs.34136		ESTs	2.6
	125054	Hs.268601	T80622	ESTs; Weakly similar to (defline not ava	2.6
	114212	Hs.21201	Z39338	ESTs; Highly similar to (defline not ava	2.6
	116959	Hs.40022	H79310	EST	2.6
45	109228	Hs.306995	AA193366	ESTs	2.6
	133989	Hs.78202	U29175	SWI/SNF related; matrix associated; acti	2.6
				Caldesmon 1, Alt. Splice 3, Non-Muscle	2.6
		Hs.285996		ESTs	2.6
		Hs.6540	Z40861	ESTs	2.6
50		Hs.171391		C-terminal binding protein 2	2.5
50		Hs.60772	AA017258	EST	2.5
	100134		D13264	macrophage scavenger receptor 1	2.5
				GA-binding protein transcription factor;	2.5
	133969	ПS./O	U13044		2.5
55		Hs.74316	AA455001	ESTS	
55		Hs.291701		oc39a08.s1 NCI_CGAP_GCB1 Homo sapiens cD	2.5
		Hs.203961		ESTs	2.5
		Hs.44583		EST	2.5
		Hs.109654	N91279	ESTs	2.5
		Hs.2785	Z19574	keratin 17	2.5
60	132883	Hs.5897	AA047151	ESTs	2.5
	102009	Hs.82643	U02680	protein tyrosine kinase 9	2.5
		Hs.20159	AA454156	ESTs	2.5
		Hs.193784		ESTs	2.5
		Hs.24908	AA256042	ESTs	2.5
65		Hs.75275	D50916	homolog of yeast (S. cerevisiae) ufd2	2.5
00		Hs.121524		glutathione reductase	2.5
		Hs.6166	AA047616	ESTs	2.5
				uridine monophosphate synthetase (orotat	2.5
		Hs.2057	AA128100	5;10-methenyttetrahydrofolate synthetase	2.5
	129045	Hs.118131	M0350	2, to monetification and or manage	2.3

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	126399	Hs.83883	AA128075	zl16d08.r1 Soares_pregnant_uterus_NbHPU	2.5
	134069	Hs.78935	U29607	Homo sapiens eIF-2-associated p67 homolo	2.5
	109816	Hs.61960	F11013	ESTs; Weakly similar to KIAA0176 [H.sapi	2.5
	134801	Hs.89695	X02160	insulin receptor	2.5
5	104232	Hs.10587	AB002351	Human mRNA for KIAA0353 gene; partial cd	2.5
	107361	Hs.159486	U72513	Human RPL13-2 pseudogene mRNA; complete	2.5
	106057	Hs.289074	AA417067	ESTs	2.5
	134252	Hs.80720	AA031782	Homo sapiens mRNA; cDNA DKFZp586B1722 (f	2.5
	128062	Hs.105547	AA379500	ESTs	2.5
10	110009	Hs.6614	H10933	ESTs	2.5
	111375	Hs.20432	N93696	ESTs	2.5
	122642	Hs.99361	AA454186	ESTs	2.5
	127999	Hs.69851	AA837495	ESTs; Weakly similar to Wiskott-Aldrich	2.5
	105029	Hs.13268	AA126855	ESTs	2.5
15	105082	Hs.26765	AA143763	ESTs: Weakly similar to Similarity to S.	2.5

TABLE 1A show the accession numbers for those primekeys lacking unigeneID's for Table 1. For each probeset we have listed the gene cluster number from which the oligonucleotides were designed. Gene clusters were compiled using sequences derived from Genbank ESTs and mRNAs. These sequences were clustered based on sequence similarity using Clustering and Alignment Tools (DoubleTwist, Oakland California). The Genbank accession numbers for sequences comprising each cluster are listed in the "Accession" column.

5

10	Pkey: CAT nu		Inique Eos probeset identifier number Bene cluster number
	Accessi		Renbank accession numbers
15	Pkey	CAT number	Accessions
		111555_1	AA071210 AA069899 AA071438 AA084912 AA084803 AA079371 AA079370
20		1596090_1	H57661 H58881
20		1606216_1	H75681 H70975
		32479_1	AB010994 U59748 AA064660
		481587	S81578
		1562851_1	H10543 R11878
	125596	1708455_1	R25698 R56582 R56018
25 30	118417	37186_1	AF080229 AF080231 AF080230 AF080232 AF080233 AF080234 BE550633 Al636743 AW614951 BE467547 Al680833 Al633818 N29986 U87592 U87593 U87590 U87591 S46404 U87587 AA463992 AW206802 Al970376 Al583718 Al672574 N25695 AW665466 Al818326 AA126128 Al480345 AW013827 AA248638 Al2144968 AA204735 AA207155 AA206262 AA204833 AW003247 AW496808 Al080480 Al631703 Al651023 Al867418 AW818140 AA502500 Al206199 Al671282 Al352545 BE501030 Al652535 BE465762 AA206331 AW451866 AA471088 AA206342 AA204834 AA206100 AW021661 AA332922 N66048 AA703396 H92278 AW139734 H92683 U87589 U87595 H69001 U87594 BE466420 Al624817
50			BE466611 Al206344 AA574397 AA348354 Al493192
	400004	007007 4	AA491830 R50173 R55192 R50320 AI732306 AI732305 AI820727 AI820728 R55191 R50319 R50227
		327827_1	
		1583542_1	H41694 H45213
25		1766315_1	R98091 W92898
35		227560_1	AA364195 AA325029 AW962050
		112052_1	AA070545 AA131490 AA131373
		231687_1	AA330501 AA661567
		232391_1	AA331503 AA332751 AW962542
40		1541209_1	T16245 R19694 F13545 H10299 T66048 T65279 H18006
40		37938_1	AF116622 AI114507 AA640834 AA377999
		112618_1	AA130614 AA071410 · · · · · · · · · · · · · · · · · · ·
		502608_1	AA906093 AA971000
	104602	524482_2	H47610 R86920
	128152	297868_1	F07973 R20353 AA442660
45	128422	1811283_1	T77794 T85681
	127897	446527_1	AA773681 AA773857
	106566	120358_1	BE298210 Al672315 AW086489 BE298417 AA455921 AA902537 BE327124 R14963 AA085210 AW274273 Al333584
		_	AI369742 AI039658 AI885095 AI476470 AI287650 AI885299 AI985381 AW592624 AW340136 AI266556 AA456390
			Al310815 AA484951
50	129735	44573_2	AJ950087 N70208 R97040 N36809 AJ308119 AW967677 N35320 AJ251473 H59397 AW971573 R97278 W01059 AW967671 AA908598 AA251875 AJ820501 AJ820532 W87891 T85904 U71456 T82391 BE328571 T75102 R34725
			AA884922 BE328517 Al219788 AA884444 N92578 F13493 AA927794 Al560251 AW874068 AL134043 AW235363
			AA663345 AW008282 AA488964 AA283144 AI890387 AI950344 AI741346 AI689062 AA282915 AW102898 AI872193
~-			AI763273 AW173586 AW150329 AI653832 AI762688 AA988777 AA488892 AI356394 AW103813 AI539642 AA642789
55			AA856975 AW505512 AI961530 AW629970 BE612881 AW276997 AW513601 AW512843 AA044209 AW856538
	•		AA180009 AA337499 AW961101 AA251669 AA251874 AI819225 AW205862 AI683338 AI858509 AW276905 AI633006
			AA972584 AA908741 AW072629 AW513996 AA293273 AA969759 N75628 N22388 H84729 H60052 T92487 AI022058
			AA780419 AA551005 W80701 AW613456 AI373032 AI564269 F00531 H83468 W37181 W78802 R66056 AI002839
			R67840 AA300207 AW959581 T63226 F04005
60	123147	2198022	AA487961
		158447_1	AA178953 AA192740
		genbank AA608	
		genbank AA180	
		tigr_HT4163	\$67998
65		tigr_HT4515	U10072
O)	100000	nAi_L114919	010072

	123798	579959_1	AA620411 AA287491
	102116	entrez_U13706	U13706
	102398	entrez_U42359	U42359
	102764	entrez_U82310	U82310
5	118475	genbank_N66845	N66845
	104776	genbank_AA026349	AA026349
	104787	genbank_AA027317	AA027317
	113702	genbank_T97307	T97307
	113938	genbank_W81598	W81598
10	122635	genbank_AA454085	AA454085
	108407	genbank_AA075519	AA075519
	108432	g	AA076626
	108555	genbank_AA084963	AA084963
	101349	entrez_L77559	L77559
15	124447	genbank_N48000	N48000
	119071	genbank_R31180	R31180
	103520	entrez_Y10511	Y10511
	103663	genbank_Z78291	Z78291
••	128046	877605_1	AA873285 Al025762
20	126959	546044_1	AA199853 AA206355
	123465	genbank AA599033	AA599033

MISSING AT THE TIME OF PUBLICATION

TABLE 2: shows a preferred subset of the Accession numbers for genes found in Table 1 which are differentially expressed in prostate tumor tissue compared to normal prostate tissue.

5 Unique Eos probeset identifier number Pkev: ExAcon: Exemplar Accession number, Genbank accession number UnigenelD: Unigene number 10 Unigene Title: Unigene gene title Ratio of tumor to normal body tissue (Relaxed ratio (87/70) UnigeneiD Unigene Title R1 ExAccn Pkey 15 37.2 131919 AA121266 Hs.272458 ESTs 120328 AA196979 Hs.290905 ESTs; Weakly similar to (defline not ava 32.6 25.2 101486 M24902 Hs.1852 acid phosphatase; prostate Hs.279477 ESTs 24.8 119073 R32894 20 Hs.183752 microseminoprotein; beta-23.8 133428 M34376 Hs.171995 kallikrein 3; (prostate specific antigen 21.4 128180 AA595348 104080 AA402971 Hs.57771 Homo sapiens mRNA for serine protease (T 18.9 Hs.162859 18.6 127537 AA569531 **ESTs** 17.4 131665 R22139 Hs.30343 **ESTs** 25 101050 K01911 Hs.1832 17.3 neuropeptide Y folate hydrolase (prostate-specific memb 17 Hs.1915 130771 N48056 16.7 107485 W63793 Hs.262476 S-adenosylmethionine decarboxylase 1 16.5 106155 AA425309 Hs.33287 **ESTs** Hs.11260 **FSTs** 129534 R73640 30 Antigen, Prostate Specific, Alt. Splice 16 100569 HG2261-HT2351 Hs.181350 kallikrein 2; prostatic 15.4 101889 S39329 Hs.99872 fetal Alzheimer antigen 15 135389 U05237 12.5 133944 AA045870 Hs.7780 **ESTs** 130974 X57985 Hs.2178 H2B histone family; member Q 11.8 35 Hs.182339 ESTs 11.8 114768 AA149007 Hs.14846 11.4 **ESTs** 104660 AA007160 ESTs; Moderately similar to KIAA0273 [H. 10.9 131061 N64328 Hs.268744 126645 Al167942 Hs.61635 Homo sapiens BAC clone RG041D11 from 7q2 10.7 Homo sapiens mRNA for JM27 protein; comp 10.6 Hs.95420 135153 N40141 40 10.6 107033 AA599629 Hs.113314 **ESTs** 118417 N66048 ESTs: Weakly similar to polymerase [H.sa 10.5 Hs.293960 10.2 126758 W37145 **ESTs** 10.1 107102 AA609723 Hs.30652 **ESTs** 116787 H28581 Hs.15641 **ESTs** 10.1 45 10 Hs.59622 115719 AA416997 **ESTs** 9.9 123209 AA489711 Hs.203270 **ESTs** 101664 M60752 Hs.121017 H2A histone family; member A 9.8 9.7 Hs.83883 112971 T17185 **ESTs** 9.7 117984 N51919 Hs.106778 ESTs 50 129523 M30894 Hs.274509 T-cell receptor; gamma cluster 9.4 132964 AA031360 Hs.167133 ESTs 9.2 121853 AA425887 Hs.98502 **ESTs** 9 119617 W47380 Hs.55999 **ESTs** 8.9 105627 AA281245 Hs.23317 8.8 **FSTs** 55 8.7 101461 M22430 Hs.76422 phospholipase A2; group IIA (platelets; 124526 N62096 Hs.293185 yz61c5.s1 Soares_multiple_sclerosis_2NbH 133845 T68510 8.2 Hs.76704 **ESTs** Hs.334762 ESTs; Weakly similar to KIAA0319 [H.sapi 133354 AA055552 8.1 119018 N95796 Hs.278695 ESTs 8 60 100394 D84276 Hs.66052 8 CD38 antigen (p45) 7.6 106579 AA456135 Hs.23023 **ESTs** 114965 AA250737 Hs.72472 7.4 **ESTs** 112033 R43162 7.1 Hs.22627 **ESTs** Human N33 protein form 1 (N33) gene, exo 102398 U42359 7 65 101201 L22524 matrix metalloproteinase 7 (matrilysin; 6.9 101803 M86546 Hs.155691 pre-B-cell leukemia transcription factor 6.8

6.8

Hs.302267 ESTs; Weakly similar to W01A6.c [C.elega

120562 AA280036

	109112	AA169379	Hs.257924	ESTs	6.8
		F10707	Hs.326416		6.7
		X07730		kallikrein 3; (prostate specific antigen	6.6
		AA219134	Hs.26691	ESTs	6.6
5		AA490969	Hs.59838	ESTs	6.6
5		U07919	Hs.75746	aldehyde dehydrogenase 6	6.5
		Z41050	Hs.108787		
		AA010163	Hs.3383	upstream regulatory element binding prot	6.5
		X07290		Human HF.12 gene mRNA	6.3
10		AA421714		Homo sapiens mRNA for KIAA0896 protein;	6.3
10		AA599267		ESTs; Weakly similar to ANKYRIN; BRAIN V	6.3
		AB000584		prostate differentiation factor	6.3
		AA609710		ESTs; Weakly similar to similar to GTP-b	6.2
		L29008	Hs.878		6.2
15		AA011176	Hs.37744	sorbitol dehydrogenase ESTs	6.2
13		AA325029	N5.01144	EST27953 Cerebellum II Homo sapiens cDN/	
			Hs.222399		6.1
		AA256485 AA053400	Hs.203213		5.9
		AA281793	Hs.72988	ESTs	5.8
20		AA491457	Hs.48948	ESTs	5.7
40		Z38839		ESTs; Weakly similar to IIII ALU SUBFAMI	5.6
		AA443993	Hs.289072		5.6
		R41933		ESTs; Weakly similar to neuronal thread	5.6
		M83822	Hs.62354	Human beige-like protein (BGL) mRNA; par	5.5
25		N41002	Hs.45107	ESTs	5.5
23		AA431407	Hs.98732	Homo sapiens Chromosome 16 BAC clone C	
		AA088851		S-adenosylmethionine decarboxylase 1	5.5
		W81598	110.202470	ESTs	5.4
		AA047036	Hs.246315		5.4
30		AA056482	Hs.7780	ESTs	5.3
50		N25110		Human guanine nucleotide exchange factor	5.3
		AA365031	Hs.98944	ESTs	5.3
		N31952		ESTs; Weakly similar to (defline not ava	5.3
		H70627		ESTs; Weakly similar to !!!! ALU SUBFAMI	5.3
35		AA284143		Homo sapiens chromosome 1 atrophin-1 rel	5.3
		W26769		ESTs; Highly similar to (defline not ava	5.2
		M21389		keratin 5 (epidermolysis bullosa simplex	5.1
		AA464728		ESTs; Weakly similar to IIII ALU SUBFAMI	5.1
		AA402613	Hs.169119		5.1
40		X91868	Hs.54416	sine oculis homeobox (Drosophila) homolo	5.1
		AA400271		ESTs; Highly similar to (defline not ava	5.1
		AA479362	Hs.47144	ESTs	5
		X07696	Hs.80342	keratin 15	5
		X52541		early growth response 1	5
45		N93839	Hs.39288	ESTs; Weakly similar to IIII ALU SUBFAMI	5

5

TABLE 2A shows the accession numbers for those primekeys lacking unigeneID's for Table 2. For each probeset we have listed the gene cluster number from which the oligonucleotides were designed. Gene clusters were compiled using sequences derived from Genbank ESTs and mRNAs. These sequences were clustered based on sequence similarity using Clustering and Alignment Tools (DoubleTwist, Oakland California). The Genbank accession numbers for sequences comprising each cluster are listed in the "Accession" column.

10	Pkey: CAT numb Accession	ber: Ge	ique Eos probeset identifier number ne cluster number nbank accession numbers
15	Pkey	CAT number	Accession
20	118417	37186_1	AF080229 AF080231 AF080230 AF080232 AF080233 AF080234 BE550633 Al636743 AW614951 BE467547 Al680833 Al633818 N29986 U87592 U87593 U87590 U87591 S46404 U87587 AA463992 AW206802 Al970376 Al583718 Al672574 N25695 AW665466 Al818326 AA126128 Al480345 AW013827 AA248638 Al214968 AA204735 AA207155 AA206262 AA204833 AW003247 AW496808 Al080480 Al631703 Al651023 Al867418 AW818140 AA502500 Al206199 Al671282 Al352545 BE501030 Al652535 BE465762 AA206331 AW451866 AA471088 AA206342 AA204834 AA206100 AW021661 AA332922 N66048 AA703396 H92278 AW139734 H92683 U87589 U87595 H69001 U87594 BE466420 Al624817
25	127248 107033 102398 113938	227560_1 235652_1 entrez_U42359 genbank_W815	

TABLE 3: shows genes, including expression sequence tags, differentially expressed in prostate tumor tissue compared to normal tissue as analyzed using the Affymetrix/Eos Hu02 GeneChip array. Shown are the relative amounts of each gene expressed in prostate tumor samples and various normal tissue samples showing the highest expression of the gene.

10	Pkey: ExAcon: Unigene Unigene R1:	ID:	Exemplar A Unigene nu Unigene ge		•	
15				·		
	Pkey	ExAccn	UnigenelD	Unigene Title	R1	
20	100235 100570 100819	D12485 D29954 HG2261-HT23 HG4020-HT42	90	phosphodiesterase l/nucleotide pyrophosp KIAA0056 protein Hs.171995 Hs.2387	6.3 5.1 Antigen, Prostate Specific, Alt. Splice Transglutaminase 10.5	9
25	101247 101416 101447 101485	L00354 L33801 M17254 M21305 M24736 M28214	Hs.89546	cholecystokinin glycogen synthase kinase 3 beta v-ets avian erythroblastosis virus E26 o Human alpha satellite and satellite 3 ju selectin E (endothelial adhesion molecul RAB3B; member RAS oncogene family	8.5 4.7 4.7 11 9.8 6.2	
30	101626 101663 101758	M57399 M60750 M77836 M81118	Hs.44 Hs.2178 Hs.79217 Hs.78989	pleiotrophin (heparin binding growth fac H2B histone family; member A pyrroline-5-carboxylate reductase 1	8.4 4.9 5.4 7.5	
35	101817 101888 102031 102052 102221	M88163 M99701 U04898 U07559 U24576	Hs.152292 Hs.95243 Hs.2156 Hs.505 Hs.3844	SWI/SNF related; matrix associated; acti transcription elongation factor A (SII)- RAR-related orphan receptor A ISL1 transcription factor; LIM/homeodoma LIM domain only 4	5.5 5.7 13.2 8.9 5.6	
40	102302 102348 102457 102473	U26173 U33052 U37519 U48807 U49957		nuclear factor, interleukin 3 regulated protein kinase C-like 2 aldehyde dehydrogenase 8 dual specificity phosphatase 4 LIM domain-containing preferred transloc	7.4 8.2 5.9 5.1 5.7	
45	102698 102751 102823 102869	U71207 U75272 U80034 U90914 X02544 X54667	Hs.29279 Hs.1867 Hs.68583 Hs.5057 Hs.572 Hs.123114	eyes absent (Drosophila) homolog 2 progastricsin (pepsinogen C) mitochondrial intermediate peptidase carboxypeptidase D orosomucolid 1	10.6 15.6 4.9 22.6 4.7	
50	103043 103093 103376 103401	X55733 X60708 X92098 X95240	Hs.93379 Hs.44926 Hs.323378 Hs.54431	eukaryotic translation initiation factor dipeptidylpeptidase IV (CD26; adenosine coated vesicle membrane protein specific granule protein (28 kDa); cyste	4.9 5.8 ⁻ 5.2 7.4	
55	103677 103962 104084 104257	Z46629 Z83806 AA298180 AA410529 AF006265	Hs.2316 Hs.83243 Hs.30732 Hs.9222	SRY (sex-determining region Y)-box 9 (ca H.saplens mRNA for exonemal dynein heavy ESTs ESTs estrogen receptor-binding fragment-assoc	5.2 4.9 6 6.4 6.8	
60	104769 104851 104896	D45332 AA025887 AA040882 AA054228 AA074880	Hs.6783 Hs.293943 Hs.10290 Hs.23165 Hs.20509	ESTs ESTs; Weakly similar to IIII ALU SUBFAMI U5 snRNP-specific 40 kDa protein (hPrp8- ESTs ESTs; Weakly similar to hypothetical pro	10.5 6.3 4.9 5.8 6.4	
65	104957 104967 105099	AA074919 AA084506 AA150776 AA233459	Hs.10026 Hs.291000	ESTs; Weakly similar to ORF YJL063c (S.c	4.8 6.5 7 5.1	

					4 ~
		AA233553	Hs.190325		4.7
	105370	AA236476	Hs.22791	ESTs; Weakly similar to transmembrane pr	10.3
		AA251330	Hs.28248	ESTs	5
				ESTs; Weakly similar to heat shock prote	8.8
5					5.5
5		AA281251		ESTs; Weakly similar to putative zinc fi	
		AA2B1623	Hs.6685	ESTs; Weakly similar to KIAA0742 protein	8
	105645	AA282138	Hs.11325	ESTs ·	14
	105691	AA287097	Hs.289068	transcription factor 4	6.3
		AA292701	Hs.5364	DKFZP5641052 protein	4.9
10				and the second s	7
10		AA393808		KIAA0438 gene product	
	105826	AA398243		ESTs; Moderately similar to similar to N	5
	105903	AA401433	Hs.200016	ESTs; Weakly similar to diphosphoinosito	9.9
	105906	AA401633	Hs.22380	ESTs	11.5
			Hs.25206	ESTs	5.1
15		AA419461	Hs.23317	ESTs	10.9
13					
		AA425367	Hs.34892	ESTs	6.6
		AA426643	Hs.10762	ESTs	8.5
	106211	AA428240	Hs.126083	ESTs	8.4
	106213	AA428258	Hs.8769	Homo sapiens mRNA; cDNA DKFZp564E153 (fr	5.7
20		AA432074	Hs.323099		5.8
20		AA443828	Hs.288856		6.3
					5.4
		AA447621	Hs.94109	ESTs	
		AA450212	Hs.42484	Homo sapiens mRNA; cDNA DKFZp564C053 (fr	
	106507	AA452584	Hs.267819	protein phosphatase 1; regulatory (inhib	5.6
25	106523	AA453441	Hs.31511	ESTs	4.7
		AA453628	Hs.37443	ESTs	4.7
		AA455087	Hs.22247	ESTs	5.7
	-				7.2
		AA456039	Hs.105421	EOT : Marks similarle Cimilarle with	
20		AA459249	Hs.8715	ESTs; Weakly similar to Similarity with	5.6
30	106820	AA481037	Hs.12592	ESTs	5.4
	106846	AA485223	Hs.34892	ESTs	5.3
	106973	AA505141	Hs.11923	Human DNA sequence from clone 167A19 on	7.5
		AA609952	Hs.12784	KIAA0293 protein	6.1
		AA620504	Hs.179898		7.1
25				ESTs; Weakly similar to ORF YKR081c [S.c	5.2
35		AA621340	Hs.10600		
		D51095	Hs.35861	DKFZP586E1621 protein	15.1
	107365	U78294	Hs.111256	arachidonate 15-lipoxygenase; second typ	4.7
	107630	AA007218	Hs.60178	ESTs	5.3
	107734	AA016225	Hs.7517	ESTs	4.8
40		AA018042	Hs.252085		7.6
-10		AA037388	Hs.82223	Human DNA sequence from clone 141H5 on c	10.5
		AA039616	Hs.173334		6.5
		AA084138	Hs.46786	ESTs	7.9
	108583	AA088276	Hs.68826	ESTs	5.6
45	108613	AA100967	Hs.69165	ESTs	6
		AA113349	Hs.69588	EST	6.3
		AA115629	Hs.118531		5.9
				ESTs; Weakly similar to PROTEIN PHOSPHAT	
		AA129968	Hs.49376		
60		AA136590		ESTs	5
50		AA147224	Hs.337232		12.7
	108948	AA149579	Hs.118258	ESTs	6.8
		AA156790	Hs.262036	ESTs	15.3
		AA171529	Hs.183887	ESTs	6.1
		AA176438	Hs.41295	ESTs	5.1
55				ESTs	5.5
22		AA196332	Hs.86043		
		AA213620		Homo sapiens mRNA; cDNA DKFZp586M1418 (
	109562	F01811	Hs.187931	ESTs; Moderately similar to voltage-gate	10.8
	109565	F01930	Hs.23648	ESTs	7
		F04600	Hs.7154	ESTs	9.9
60		F10770		Homo sapiens clone 669 unknown mRNA; com	6.4
00				ESTs	5.3
		H02308	Hs.20792		
		H20276	Hs.31742	ESTs	16.8
•	110854	N32919	Hs.27931	ESTs	10
	110924	N47938	Hs.12940	yy84a09.s1 Soares_multiple_sclerosis_2Nb	5.6
65		N55514	Hs.318584		6.9
		N59858	Hs.33032		
		N66613	Hs.99364	ESTs	5
			Ho 400400	ESTs; Weakly similar to IIII ALU CLASS C	5.6
		N66857			
	111221	N68869	Hs.15119	ESTs	6.2

	111348		Hs.9585	ESTs	5.4
	111353	N90430	Hs.6616		5.3
	111495	R07210	Hs.9683	ESTs	5.8
	111540	R08850	Hs.9786	ESTs	6
5	111579	R10657	Hs.167115	KIAA0830 protein	12.6
-	111581	R10684	Hs.5794	ESTs	7.1
		R25375	Hs,128749	ESTs	6.2
		R37460	Hs.25231	ESTs	9.4
		R37778	Hs.18685	ESTs; Weakly similar to hypothetical pro	6.5
10		R40431	Hs.14846	Homo sapiens mRNA; cDNA DKFZp564D016 (fr	
10		R42036	Hs.6763	KIAA0942 protein	6.4
				•	5.6
		R49173	Hs.330242		
		R53765		KIAA0981 protein	9.3
1.5		R59740	Hs.5740	ESTs	4.7
15		R63841	Hs.157461		6
	112601	R79111	Hs.78225	annexin A1	5.4
	112753	R93696	Hs.169882	ESTs	5.8
	112902	T09262	Hs.129190		5.1
	112984	T23457	Hs.289014	ESTs	4.9
20	113021	T23855	Hs.129836	KIAA1028 protein	10.8
	113083	T40530	Hs.266957	ESTs; Weakly similar to heat shock prote	5.7
		T57773	Hs.10263	ESTs	7.3
		T88878	Hs.86538	ESTs	8.7
		W60439	Hs.8858	ESTs; Moderately similar to cbp146 [M.mu	4.9
25		W72382	Hs.11958	oxidative 3 alpha hydroxysteroid dehydro	4.7
		W85765	Hs.30504	Homo sapiens mRNA; cDNA DKFZp434E082 (fr	
		W87462	Hs.21894	ESTs	5.9
		W87544	Hs.268828		4.7
		Z38595		ESTs; Highly similar to KIAA0886 protein	21.3
30			Hs.143611		9.6
50		Z41395	Hs.130489		5.2
		Z41450		Bicaudal D (Drosophila) homolog 1	7.4
		AA018216			8.2
		AA025370	Hs.40109		5.4
25		AA101416		ESTs; Weakly similar to PTB-ASSOCIATED S	4.8
35		AA131450	Hs.103822		
		AA133527		ESTs; Weakly similar to The KIAA0138 gen	5.1
		AA234362	Hs.87159		5.5
		AA235112	Hs.42179	ESTs; Moderately similar to similar to m	6.3
40		AA235811	Hs.293672		5.2
40		AA236177	Hs.76591	KIAA0887 protein	4.7
	114908	AA236545	Hs.54973	ESTs	5.2
	114932	AA242751	Hs.16218	KIAA0903 protein	5.7
	115084	AA255566	Hs.42484	Homo sapiens mRNA; cDNA DKFZp564C053 (fr	
	115140	AA258030	Hs.279938	ESTs; Weakly similar to supported by GEN	5.9
45	115468	AA287061	Hs.48499	ESTs; Highly similar to Bdeight protein	4.7
	115583	AA398913	Hs.45231	LDOC1 protein	7.6
	115709	AA412519	Hs.58279	ESTs	4.8
	115772	AA423972	Hs.131740	ESTs	5
		AA424029	Hs.288390	ESTs; Moderately similar to dynamin; int	5.4
50	115776	AA424038	Hs.81897	ESTs	5
• •		AA427528	Hs.130965	ESTs; Weakly similar to ZINC FINGER PROT	13.7
		AA446121	Hs.44198	Homo sapiens BAC done RG054D04 from 7q3	10.6
		AA451748	Hs.83883	Human DNA sequence from clone 718J7 on c	6.8
		AA457566	Hs.28777	ESTs	6
55		AA459117	Hs.31575	SEC63; endoplasmic reticulum translocon	7.3
<i>JJ</i>		AA460701	Hs.15423	ESTs	5.5
		AA489033	Hs.62601	Homo sapiens mRNA; cDNA DKFZp586K1318 (
				ESTs	5.9
		AA521472	Hs.71252		5.9
60		AA599463		protein phosphatase 2 (formerly 2A); reg	
60		AA599963	Hs.59698	ESTs	7.9
		AA609219	Hs.39982	ESTs	9.2
		D59325	Hs.121429		5.2
		D80055	Hs.45140	ESTs	4.9
		F09156	Hs.66095	ESTs	7.2
65	116722	F13654		HSFIH32 Stratagene cat#937212 (1992) Hom	5.5
	116766	H13260	Hs.95097	ESTs	5.9
	117453	N29568	Hs.108319	thyrold hormone receptor-associated prot	6.9
	117557	N33920	Hs.44532	diubiquitin	4.8
		N45114	Hs.126280	•	6.3
	-				

	440004	NICO4E4	Un 47447	ECT	44.4
		N52151 N62339	Hs.47447	ESTs heat shock 90kD protein 1; alpha	11.4 6.2
		N69207	Hs.203697		5.8
		N70358		growth hormone receptor	7.1
5		N89881	Hs.44577	y	6
-		N94303	Hs.55028	ESTs	9.3
		R42424	Hs.63841	ESTs	6
		R45175	Hs.117183	ESTs	17.9
	119271	T16387	Hs.65328	ESTs	6
10	119367	T78324	Hs.250895		5
		W69440	Hs.48376		15.4
		W70205		kinesin family member 3A	10.1
		W72967		ESTs; Weakly similar to hypothetical pro	5.3
15 .		Z41078	Hs.66035	ESTs	4.8
15		AA173939		ESTs; Weakly similar to inner centromere ESTs; Highly similar to NY-REN-62 antige	8.8 4.9
		AA190888 AA236010	Hs.26613		
		AA253400		turnor protein 63 kDa with strong homolog	5.6
		AA261852	Hs.192905		4.9
20		AA280738	Hs.34892	ESTs	8.8
		AA282074	Hs.237323	ESTs	6.2
	120713	AA292655	Hs.96557	ESTs	9.9
		AA398246	Hs.97594	ESTs	16.4
05		AA406293	Hs.41167		6.9
25		AA412049	Hs.290347		7.6
		AA412105 AA424814	Hs.193736		5.8 4.6
		AA431302	Hs.48827 Hs.98721	EST; Weakly similar to N-copine [H.sapie	5.6
		AA437311	Hs.98927	ESTs .	5.7
30		AA446859	Hs.99083	ESTs	6.5
		AA460158		KIAA1028 protein	12.4
	122792	AA460225	Hs.99519	ESTs	5.1
		AA478539	Hs.104336		4.9
25		AA485724	Hs.27413		5.4
35		AA485957		Homo sapiens clone 25032 mRNA sequence	5 4.7
		AA495981 AA496252	Hs.250830 Hs.105069		7.A
		AA609006	Hs.111240		9.1
		AA609200	113.111240	ESTs	4.7
40		AA609310	Hs.188691		4.8
••		AA609651	Hs.112742		7
	123968	C14333	Hs.108327	damage-specific DNA binding protein 1 (1	5
		H45996	Hs.97101	putative G protein-coupled receptor	6.8
45		N21626	Hs.102406	==::	102
45		N22401	11-400070	yw37g07.s1 Morton Fetal Cochlea Homo sap	10.6
		N58172	Hs.109370 Hs.174195	•	14.2 4.8
		R88992 W38419	NS.174185	ESTS	4.7
		W01626		za36e07.r1 Soares fetal liver spleen 1NF	5.1
50		AA947601	Hs.97056	ESTs	5.1
	400040	Z36290	Hs.173933	ESTs; Weakly similar to NUCLEAR FACTOR 1	4.6
		AA662913	Hs.190173	ESTs	5
	127308	AA507628	Hs.334390		4.8
		AI024352		immunoglobulin superfamily; member 4	4.7
55		AI457411	Hs.106728		4.8
		AA828760	Hs.292059		4.8 5
		A1400862 A1039722	Hs.265130 Hs.279009		5.8
		A1039722 A1088155		ESTs; Weakly similar to unknown [H.saple	17
60		AA176446		ESTs; Weakly similar to hypothetical 43.	4.8
•		L38608	Hs.10247		7.9
		AA242816		ESTs; Weakly similar to KIAA0437 [H.sapi	8.1
	128651	AA446990	Hs.103135	ESTs	6.5
	129088	AA215971		KIAA0992 protein	5.2
65		N26391	Hs.250723		5.1
		AA234048	Hs.7753	calumenin	5.8
		AA211941		polyadenylate binding protein-interactin	5.8
		N27524 AA410311	Hs.260024 Hs.44208	Cdc42 effector protein 3 ESTs	5.2 5.1
	12040/	WHIMII	, 10. THEOU	2010	J. I

	129564	H22136	Hs.75295	quanylate cyclase 1; soluble; alpha 3	16.3
		AA458578	Hs.12017	KIAA0439 protein; homolog of yeast ubiqu	9.2
	129821		Hs.12696	cortactin SH3 domain-binding protein	8.6
_	129823	X00948	Hs.105314	relaxin 2 (H2)	9.1
5	129847	W46767	Hs.296178	ESTs; Weakly similar to RNA POLYMERASE I	5.4
_	129912	AA047344		ESTs; Highly similar to NY-REN-6 antigen	6.5
	129958		Hs.1378	annexin A3	5.1
	129977	J040/6	Hs.1395	early growth response 2 (Krox-20 (Drosop	8.6
•	130061	U82256	Hs.172851	arginase; type II	7.4
10	130241	U78313	Hs.153203	MyoD family inhibitor	4.9
		N21679	Hs.180059	1 _ ·	5.8
				neurofilament; light polypeptide (68kD)	6.7
	130541				
	130619	AA477739	Hs.12532	ESTs	6.4
	130925	N71935	Hs.169378	multiple PDZ domain protein	7.9
15	130938	AA013250	Hs.21398	ESTs; Moderately similar to PUTATIVE GLU	6.2
	130971	H20332	Hs 301444	signal sequence receptor; gamma (translo	6.4
	131066		Hs.22588	ESTs	5
					-
	131126			myotubularin related protein 2	6.4
	131310	J02960	Hs.2551	adrenergic; beta-2-; receptor; surface	7.9
20	131487	AA253220	Hs.27373	Homo sapiens mRNA; cDNA DKFZp564O1763 (5.9
	131561		Hs 294101	pre-B-cell leukemia transcription factor	7.6
		U90551	Hs.28777	H2A histone family; member L	5.1
		N62922	Hs.29088	ESTs	11
	131629	AA442119	Hs.238809		4.9
25	131682	AA428368	Hs.30654	ESTs	4.8
		R68657	Hs.90421	ESTs; Moderately similar to !!!! ALU SUB	6.5
		N32724	Hs.32317		5.6
				Sox-like transcriptional factor	
		H93381	Hs.38085	ESTs; Weakly similar to putative glycine	7.2
	132122	U65092	Hs.40403	Cbp/p300-interacting transactivator, wit	5.6
30 ·	132191	AA449431	Hs.288361	KIAA0741 gene product	8
		AA608856	Hs.431	murine leukemia viral (bmi-1) oncogene h	5.5
		AA429478		ESTs; Highly similar to CGI-49 protein [6.6
		AA021608	Hs.172510		5.8
	132572	AA448297	Hs.237825	signal recognition particle 72kD	6.2
35	132581	R42266	Hs.52256	ESTs; Weakly similar to beta-TrCP protei	16
		N47109	Hs.5521	ESTs	6.8
		AA279359	Hs.55220	BCL2-associated athanogene 2	5.3
	132725		Hs.184167		7.8
	132783	N74897	Hs.278894	DEAD/H (Asp-Glu-Ala-Asp/His) box polypep	5.9
40	132790	X75535	Hs.168670	peroxisomal famesylated protein	8
	132939	U76189	Hs.61152	exostoses (multiple)-like 2	5.2
	133142		Hs.65874	ESTs	5.2
					_
		U29589	Hs.7138	cholinergic receptor; muscarinic 3	10.3
	133434	AA278852	Hs.30212	ESTs	5.8
45	133453	M68941	Hs.73826	protein tyrosine phosphatase; non-recept	4.9
	133520	X74331	Hs.74519	primase; polypeptide 2A (58kD)	13.1
		T33873	Hs.74624	protein tyrosine phosphatase; receptor t	4.6
					4.8
		D13315	Hs.75207	glyoxalase I	
	133626	H75939	Hs.75277	Homo sapiens mRNA; cDNA DKFZp586M141 (fr	
50	133633	D21262	Hs.75337	nucleolar phosphoprotein p130	6.3
	133797	S66431	Hs.76272	retinoblastoma-binding protein 2	6
		N34096	Hs.7766	ubiquitin-conjugating enzyme E2E 1 (homo	5.4
					5.2
		U47414	Hs.79069	cyclin G2	
	134249	N89827	Hs.80667	RALBP1 associated Eps domain containing	6.5
55	134321	AA418230	Hs.8172	ESTs	7
	134453	X70683	Hs.83484	SRY (sex determining region Y)-box 4	4.7
		X57025	Hs.85112	Insulin-like growth factor 1 (somatomedi	7.7
		U66615		SWI/SNF related; matrix associated; acti	6.4
	134592	U82613	HS.289104	Alu-binding protein with zinc finger dom	5.4
60	134654	W23625	Hs.8739	ESTs; Weakly similar to ORF YGR200c [S.c	5
	134666	AA482319	Hs.8752	putative type II membrane protein	5.4
		Z49099	Hs.89718	spermine synthase	6.7
		AA431480	Hs.169358		9.8
		X04602	Hs.93913	Interleukin 6 (Interferon; beta 2)	5.7
65	135155	AA358268	Hs.166556	ESTs; Moderately similar to transcriptio	4.9
		L10333	Hs.99947	reticulon 1	5.3
		M10098	30	AFFX control: 18S ribosomal RNA	4.6
			No FFC40		
		AW079607		ESTs; Weakly similar to ZnT-3 [H.sapiens	7.8
	300273	AW013907	Hs.167531	ESTs; Moderately similar to predicted us	11.5

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	300319	AW157646	Hs.153506	ESTs; Weakly similar to microtubule-acti	8.5
	300566	H86709	Hs.326392	son of sevenless (Drosophila) homolog 1	5.8
		AI989417	Hs.134289		4.4
				ESTs	7.9
_		Al239706	Hs.93810		
5		AA039352		ESTs; Weakly similar to ORF YDL040c [S.c	4.5
	300680	AW468066	Hs.24817	ESTs; Weakly similar to KIAA0986 protein	5.2
	300762	AI497778	Hs.20509	ESTs	6.4
		AI076890	Hs.146847		5.8
					10.6
10		AA406411		ESTs; Weakly similar to KIAA0989 protein	
10		A1863068		ESTs; Weakly similar to putative zinc fi	5.6
	300834	AF109300	Hs.147924	ESTs	6.7
	300923	AW136372	Hs.1852	ESTs	7.6
		AA593373	Hs.293744		5.5
					7
1.5		AA947682	Hs.20252	ESTs; Weakly similar to Chain A; Cdc42hs	
15		AI659131	Hs.197733		24.9
	301242	AW161535	Hs.23782	ESTs	11.8
	301254	Al049624	Hs.283390	EST cluster (not in UniGene) with exon h	4.3
		H29500	Hs.7130	ESTs; Moderately similar to N-copine (H.	4.3
		AA156879		ESTs; Weakly similar to ZINC FINGER PROT	6.6
20					
20		AI802946		ESTs; Weakly similar to match to ESTs AA	5.7
	301656	AW008475	Hs.151258	EST cluster (not in UniGene) with exon h	6.8
	301689	Z44810	Hs.301789	ESTs; Weakly similar to similar to C.ele	6.3
	301783	AL046347	Hs.83937	Homo sapiens PAC clone DJ1159O04 from 7p	6.2
		A1800004		ESTs; Weakly similar to MesP1 [M.musculu	8.5
25					4.6
ZJ		R20002	Hs.6823	ESTs; Weakly similar to intrinsic factor	
	301891	AF131855		Homo sapiens clone 25056 mRNA sequence	6.3
	302005	A1869666	Hs.123119	ESTs	36.8
	302056	Al457532	Hs.30488	ESTs: Moderately similar to ROSA26AS [M.	9.5
		H05698		ESTs; Weakly similar to protein-tyrosine	5.8
30		AL021397		ribosomal protein L34 pseudogene 1	8.8
20					
		AB022660		KIAA0437 protein	5.9
	302214	AJ001454		Homo sapiens mRNA for testican-3	4.3
	302236	Al128606	Hs.6557	zinc finger protein 161	4.3
	302358	D81150	Hs.322848	EST cluster (not in UniGene) with exon h	5.5
35		NM_004917		EST cluster (not in UniGene) with exon h	26.8
JJ		AC003682		multiple UniGene matches	8.2
					6.4
		NM_000522		EST cluster (not in UniGene) with exon h	
		AA425562		EST cluster (not in UniGene) with exon h	5
	302792	AA343696	Hs.46821	ESTs; Weakly similar to putative [H.sapi	4.8
40	302881	AA508353	Hs.105314	relaxin 1 (H1)	78.8
-	302892	N58545	Hs.42346	histone deacetylase 3	8.5
		AW118352		EST cluster (not in UniGene) with exon h	7.4
				EST cluster (not in UniGene) with exon h	5.5
		AW263124	NS.313111		
4.5		AF199613		EST cluster (not in UniGene) with exon h	4.6
45	303125	AF161352		EST cluster (not in UniGene) with exon h	5.8
	303280	A1571580	Hs.170307	ESTs	4.3
	303306	AA215297	Hs.61441	EST cluster (not in UniGene) with exon h	6.4
		AL134164	Hs.145416		6.6
		AA255977			19.5
50				ESTs; Highly similar to ubiquitin-conjug	
50		AA298471		EST cluster (not in UniGene) with exon h	6.6
	303401	AA758552	Hs.309497	ESTS	6.8
	303525	AW516519	Hs.273294	ESTs	4.8
		AA348111	Hs.96900	ESTs	12.1
		AA355607		ESTs; Weakly similar to MMSET type I [H.	8.2
55			Hs.242540		8.4
23		AW338520		· · · · · · · · · · · · · · · · · · ·	
		AW500106	Hs.23643	EST cluster (not in UniGene) with exon h	4.9
	303699	D30891	Hs.19525	EST cluster (not in UniGene) with exon h	15.7
	303702	AW500748	Hs.224961	ESTs; Weakly similar to 73 kDA subunit o	6.3
		AI741397	Hs.114658		4.6
60	202700	AA521510	Hs.145010		12.5
JU	000722	AMERICA			
		AW502405		ESTs; Weakly similar to tumor suppressor	4.3
		AA707750		ESTs; Weakly similar to cis-Golgi matrix	5.4
	303752	A1017286	Hs.5957	EST cluster (not in UniGene) with exon h	5.3
		AW503733	Hs.9414	ESTs	13
65		Al275850		EST cluster (not in UniGene) with exon h	7.8
<i>U.</i>				translocase of inner mitochondrial membr	4.8
	304053	R00493			
	304218	N66373	Hs.27973		6
	305200	AA668128	Hs.45207	EST singleton (not in UniGene) with exon	5.7
	306716	AI024916	Hs.251354	ESTs	5.7

	307848	Al364186		EST singleton (not in UniGene) with exon	7.3
•		Al368665	Hs.31476	EST singleton (not in UniGene) with exon	5.4
		Al460004	Hs.31608	EST singleton (not in UniGene) with exon	8.1
5		Al613519		EST singleton (not in UniGene) with exon	5.5 4.4
3		Al863051 Al927149	Hs.279815 Hs.29797		4.5
		AW075342	Hs.9271	EST singleton (not in UniGene) with exon	7.4
		AW205604		ESTs; Weakly similar to IIII ALU SUBFAMI	5
		Al921750	Hs.144871		5
10		Al685841	Hs.161354		11.6
		AJ478629 AJ262148	Hs.158465 Hs.145569		5.8 9.7
		AI202140 AI734009		EST cluster (not in UniGene)	10.4
		AI612775	Hs.145710		4.6
15		Al420227	Hs.149358		72.9
		AW292180	Hs.156142		7.6
		Al338013	Hs.140546		9.2
		AW269082 AW262580	Hs.175162 Hs.147674		4.5 4.9
20		A1973051	Hs.224965		7.6
20		Al655662	Hs.197698		41.3
		Al767957		ESTs; Weakly similar to Y38A8.1 gene pro	4.5
•		Al679524		ESTs; Moderately similar to III! ALU SUB	4.6
25		AW136713	Hs.23862	ESTS	5.9
25		A1824863 A1828254	Hs.211420 Hs.271019		4.8 5.8
		AI682088	Hs.79375	ESTs	26.4
			Hs.27133		6.4
		AW025661	Hs.240090	ESTs	7.4
30		AI682478	Hs.13528		4.6
		AA765470 AW014013	Hs.85092 Hs.107056	ESTS	6.7 5.3
		R16890	Hs.137135		5.6
		AW451654	Hs.257482		4.3
35		AA759250		cytochrome b-561	11
		AA834800		EST cluster (not in UniGene)	16.9
		Al380207	Hs.125276		4.7
		C01367 R46180	Hs.127128 Hs.153485		5.3 6.2
40		AA847398	Hs.291997		4.8
		R49353	Hs.293892		5.2
	312480	R68651	Hs.144997		9.5
		C17785	Hs.182738		6.3
45		AA033609	Hs.239884		11.2 4.7
43		Al695522 Al004377	Hs.191271 Hs.200360		7
		Al623511	Hs.118567		5.1
		AA976064	Hs.180842		6.5
~~		AA694607		EST cluster (not in UniGene)	10.8
50		AA772279	Hs.126914		5
		A1813654 AA939266	Hs.5957 Hs.278626	ESTS	5.8 7.7
		H92571	Hs.234478		6.5
		AA836271	Hs.125830		4.6
55	312983	A1079278	Hs.269899	ESTs	5.1
		AA249018		EST cluster (not in UniGene)	7
		N36417	Hs.144928		6.3 4.3
		Al801098 Al039702	Hs.151500	collagen; type I; alpha 2	4.8
60		AA827805	Hs.124296		5
		Al200281	Hs.123910	ESTs	5.9
		Al420611	Hs.127832		4.6.
		A1088120	Hs.122329		7.4
65		AA745689 Al261390		ESTs; Weakly similar to similar to zinc	6.3 5.6
O)	313540	A1797301	Hs.146085 Hs.5740	ESTs	5.0 5.9
	313568	AW467376	Hs.129640		4.3
	313569	Al273419	Hs.135146	ESTs; Weakly similar to ZK1058.5 [C.eleg	4.6
	313603	AW468119		EST cluster (not in UniGene)	6.8

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		AW295194		DKFZP434N126 protein	5.2
	313625	AW468402	Hs.254020	ESTs	7.8
	313634	AA688292	Hs.337786	ESTs	4.4
	313635	AA507227	Hs.6390	ESTs	8.1
5	313638	A1753075	Hs.104627	ESTs	6.7
	313670	C16690	Hs.23767	EST duster (not in UniGene)	4.4
	313671	W49823	Hs.104613	ESTs	4.4
	313676	AA861697	Hs.120591	EST cluster (not in UniGene)	13.4
	313703	Al161293		ESTs; Weakly similar to KIAA0525 protein	10
10	313712	AA768553	Hs.74170	ESTs	5.2
		AW296132	Hs.55098	ESTs	5.4
	313979	AI535895	Hs.221024	ESTs	4.3
		AI732100	Hs.187619		13.6
		AW245993	Hs.223394		6.4
15		AI821895	Hs.193481	ESTs	29.4
	314188	AL138431	Hs.164243		4.6
		AL036001	Hs.48376	ESTs	5.7
		AA743396	Hs.189023	ESTs	4.9
		AA732359	Hs.96264	ESTs	4.4
20		AA731431		EST cluster (not in UniGene)	6.4
		Al280112	Hs.125232		5.3
		AI754701		ESTs; Weakly similar to afternatively sp	6.2
		AI052358	Hs.193726		4.5
		AW207206	Hs.136319		17
25	•	AW502698	Hs.118152		8.9
		AI538226	Hs.32976		9.4
		AA481027		ESTs; Weakly similar to ORF YGR245c [S.c	8
		AA493811	Hs.294068		6
		Al672225	Hs.222888	ESTs	19.3
30		AA548906	Hs.122244		4.5
		AA521381	Hs.187726		5.3
		AA524953			4.6
		AA533447		EST cluster (not in UniGene)	5.1
		AW292425	Hs.163484		15.5
35		AA876910	Hs.134427		20
		AW452948	Hs.257631		5.3
		Al821085		ESTs	8.2
		Al915927	Hs.34771		5.4
		Al420753	Hs.66731	ESTs	5.1
40		Al985544	Hs.12450	ESTs	5.8
		Al222165	Hs.144923		4.5
		AW291563	Hs.104696	ESTs	8
		AA764918	Hs.256531		4.8
		Al263393	Hs.145008	ESTs ·	6.2
45	315379	Al378329	Hs.126629	ESTs	5.4
		AW293424	Hs.75354	ESTs	5.1
	315442	AA977935	Hs.127274	ESTs	6.6
	315443	AW003416	Hs.160604	ESTs	5.5
	315528	R37257	Hs.184780	ESTs	8.1
50	315593	AW198103	Hs.158154	ESTs	9.9
	315634	AA837085	Hs.220585	ESTs	7.8
	315705	AW449285	Hs.313636		8.9
	315707	Al418055	Hs.161160	ESTs	5.1
	315714	AA744015		EST cluster (not in UniGene)	6.1
55	315740	T05558	Hs.156880	EST cluster (not in UniGene)	6.8
	315762	Al391470	Hs.158618		5.3
		AA744875	Hs.189413	ESTs	5
	315843	AA679430	Hs.191897	ESTs	5.7
		Al800041	Hs.190555	ESTs	9.2
60	316012	AA764950	Hs.119898	ESTs	4.3
		AA708016	Hs.190389	ESTs	5.9
		AA693880		EST cluster (not in UniGene)	6.7
	316074	AW517542	Hs.293273		5.5
		AW203986	Hs.213003		5.1
65		Al127483	Hs.120451		8.2
-		AA760894	Hs.153023	ESTs	17.1
		AA768025	Hs.186854	EST	4.6
		AW135854	Hs.132458	ESTs	4.3
		AW015940	Hs.232234		7.6

		AA831215			5.1
		AW138241	Hs.210846		6.4
		AW051597	Hs.143707		4.4
~		AA864968	Hs.127699		11
5		AW445167	Hs.126036		13.5
		D56760	Hs.93029 Hs.126594	= 11	8.7 8.7
		AI806867	Hs.120394		6.7 11.1
		AA931245 Al654187	Hs.195704		14.2
10		AW292779	Hs.169799		5.8
10		AI733277	Hs.128321		5.4
		N29974		EST duster (not in UniGene)	11.4
		AW295184	Hs.129142	ESTs; Weakly similar to DEOXYRIBONUCLEAS	
		A1828602	Hs.211265		5.3
15		AI565071	Hs.159983	ESTs	7.7
	318239	AI085198	Hs.164226	ESTs	13.1
	318268	AJ817736	Hs.182490	ESTs	6.2
•		AW294013	Hs.200942		4.6
20		R45530	Hs.1440	gamma-aminobutyric acid (GABA) A recepto	6
20		Al949409	Hs.194591		12.3
		Al151010	Hs.157774		4.3
		AW291511	Hs.159066	EST duster (not in UniGene)	25.9 7
		T30280 AW206806	Hs.115325		4.8
25		A1133617	Hs.10177		5.5
 _		AW175665	Hs.278695		5.7
		Al493742	Hs.165210		11
		W26276	Hs.136075		5.9
	318753	AA578265	Hs.7130	copine IV	5.5
30	319080	Z45131	Hs.23023	ESTs	16.9
		F06504	Hs.27384		4.6
		AF071538	Hs.79414	prostate epithelium-specific Ets transcr	6.6
		R21054	Hs.180532		4.9
35		D78808	Hs.283683		8.2 9.3
33		AA621606 AA460775	Hs.117956 Hs.6295	ESTS	14.3
		AA424266		EST cluster (not in UniGene)	12.8
		AA337642		nuclear factor related to kappa B bindin	5.1
		AA179304		ESTs; Moderately similar to !!!! ALU SUB	4.3
40		T80579	Hs.290270		5.8
	320076	AI653733	Hs.271593		8.5
		AW296219		RAB7; member RAS oncogene family-like 1	9.8
		T99949		EST cluster (not in UniGene)	9.8
45		AL039402		DEME-6 protein	7.9
45	-	AF071202		ATP-binding cassette; sub-family C (CFTR	56.2 8.3
		R49889 Al089817	Hs.24144 Hs.237146		5.4
		NM_006953		EST cluster (not in UniGene)	7
		AL049443	Hs.161283	Homo sapiens mRNA; cDNA DKFZp586N2020 (f	
50		AL049977	Hs.162209	Homo sapiens mRNA; cDNA DKFZp564C122 (fr	6.7
•		AW263086	Hs.118112		6
	320796	AF038966	Hs.31218	secretory carrier membrane protein 1	13.
	320800	A1681006	Hs.71721	ESTs	6.2
		AW360847	Hs.16578		9.3
55		AI473796	Hs.135904	ESTs	8.1
		D59945	Hs.65366		6
		AA633772	Hs.116796	- 1 .1	9.2 5
		AW195012 H19732	Hs.293970 Hs.247917		5.9
60		AA018386	Hs.64341	ESTs	4.6
UU		H52462		EST cluster (not in UniGene)	5.8
		AB033041		EST cluster (not in UniGene)	8.4
		AW372449		EST duster (not in UniGene)	7.3
		AW297633	Hs.118498		14.
65		H80483	Hs.46903	EST duster (not in UniGene)	9.2
	321609	H86021		ESTs; Weakly similar to hMmTRA1b [H.sapl	4.8
		Al791838	Hs.193465		5.5
		Al356352	Hs.108932		4.6
	321644	Al204177	Hs.237396	2018	6.6

	321681	AA233821	Hs.190173	EST cluster (not in UniGene)	4.6
	321726	X91221		EST cluster (not in UniGene)	5
	321758	U29112		EST cluster (not in UniGene)	6.2
	321877	AL109784	Hs.189222	EST cluster (not in UniGene)	4.6
5	321899	N55158	Hs.29468	ESTs	4.6
	321902	AA746374	Hs.145010	ESTs	8.2
	322007	AW410646	Hs.164649	ESTs	5.1
	322055	AL137646	Hs.146001	EST cluster (not in UniGene)	4.3
	322092	AF085833	Hs.135624	EST cluster (not in UniGene)	4.3
10	322221	AI890619	Hs.179662	nucleosome assembly protein 1-like 1	4.4
	322278	AF086283		EST cluster (not in UniGene)	5.8
	322303	W07459	Hs.157601	EST cluster (not in UniGene)	22
	322437	AW393804	Hs.170253	ESTs; Weakly similar to rabaptin-4 [H.sa	4.4
	322493	AF143235	Hs.279819	EST cluster (not in UniGene)	7.2
15	322782	AA056060	Hs.202577	EST cluster (not in UniGene)	18.4
	322811	AA782292	Hs.105872	ESTs	6.9
	322818	AW043782	Hs.293616	ESTs	10.7
	322826	Al807883	Hs.180059		5
	322887	Al986306	Hs.86149	ESTs; Weakly similar to KIAA0969 protein	11.9
20	322889	AA081924	Hs.124918		7.1
	322924	AA669253	Hs.136075		4.5
	322982	Al351191	Hs.128430		6.6
		AA422116	Hs.191461		4.7
		AA336609	Hs.10862	ESTs	6.9
25		AL118747	Hs.26691	EST cluster (not in UniGene)	8.3
		AA148950	Hs.188836		4.6
		AL118923		EST cluster (not in UniGene)	7.5
		AA157726	Hs.264330		7.5
20		AA157867	Hs.5722	ESTs	4.7
30		Z44354		guanine nucleotide binding protein (G pr	4.9
		AA176982		EST cluster (not in UniGene)	6.1
		AL120351		EST cluster (not in UniGene)	4.3
		Al827137	Hs.336454		6.2
35		AF131846		Homo sapiens clone 25028 mRNA sequence	6.3
33		AF055019	Hs.21906	·	12.6
		AA363148 AI829770	Hs.293960 Hs.190642		10.9 7.6
		AA836452	Hs.323822		7.6
		AA639902	Hs.104215		24.7
40		A1655499	Hs.161712		14.1
••		AL134875	Hs.108646		5.3
		AL135067	Hs.117182		6.1
		C05278		ESTs; Moderately similar to [PYRUVATE DE	8.5
	323496	AI826801	Hs.300700		4.5
45	323507	H71721	Hs.128387	ESTs	4.4
	323545	AI814405	Hs.224569		5.8
	323623	AA314280	Hs.146589	EST cluster (not in UniGene)	5
	323663	AW263526	Hs.243023		7.7
	323691	AA317561	Hs.145599	EST cluster (not in UniGene)	5.9
50	323810	AA740405	Hs.108806	ESTs	6.2
	323846	AA337621	Hs.137635	ESTs	6
		AA354940	Hs.145958		10.7
		AI636775	Hs.6831	ESTs	5.4
ہے ہے		AA367032	Hs.217882		5.8
55		AA844907	Hs.274454	EST cluster (not in UniGene)	4.4
		AW177009		EST cluster (not in UniGene)	4.6
		AL046575	Hs.130198		11
		Al146686	Hs.143691		13.7
<i>4</i> 0		Al524039	Hs.192524		6.8
60		AA627642	Hs.4994	transducer of ERBB2; 2 (TOB2)	4.9
		AA884766	U- 0040.~	EST duster (not in UniGene)	4.3
		F28212		EST cluster (not in UniGene)	4.7
		AA464018		EST duster (not in UniGene)	13.6
65		AW014022	Hs.170953		7.6
UJ		AW501974	Hs.74170		5.6 24.2
		AW016378	Hs.292934 Hs.195839		24.2 54
		AA508552 Al346282	Hs.87159		4.6
		AA448021	Hs.94109		5.7
	GE TOEU		. 10.0 7 100	and the state of t	J.1

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9
        324626 Al685464
                                       ESTs
        324658 Al694767
                                                                                22
                            Hs.129179 ESTs
                                                                                 4.9
                            Hs.112451 ESTs
        324676 AW503943
                            Hs.293341 ESTs; Weakly similar to Pro-a2(XI) [H.sa
                                                                                 10.6
        324691 Al217963
 5
                            Hs.257339 ESTs
                                                                                 10.2
        324696 AA641092
        324713 AW340249
                            Hs.163440 ESTs
                                                                                 5.5
                            Hs.131798 EST cluster (not in UniGene)
                                                                                 7.2
        324715 AI739168
                                                                                 34.4
                            Hs.116467 ESTs
        324718 Al557019
        324720 AA578904
                            Hs.292437 ESTs
                                                                                 4.8
10
                                                                                 7.9
                            Hs.272072 ESTs; Moderately similar to IIII ALU SUB
        324752 Al279919
                            Hs.144871 EST cluster (not in UniGene)
                                                                                 5.2
        324753 AA612626
        324790 Al334367
                             Hs.159337 ESTs
                                                                                 7.6
                                                                                 12.6
                            Hs.14553 ESTs
        324801 Al819924
                                                                                 6.5
        324804 Al692552
                                       FSTs
15
                            Hs.337533 ESTs
                                                                                 4.5
        324845 AA361016
                            Hs.136102 KIAA0853 protein
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        324888 Al564134
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        324929 AI741633
                            Hs.125350 ESTs
                                       EST cluster (not in UniGene)
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        324961 AA613792
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                            Hs.22380
                                       ESTs
        325108 AA401863
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                                       CH.20_hs gi[6552458
                                                                                 9.6
        326816
                                       CH.21_hs gij5867660
                                                                                 4.8
        326997
                                       CH.21_hs gi|6682516
                                                                                 4.3
        327098
                                       CH.07_hs gij5868455
                                                                                 5.8
        328492
                                       CH.X_hs gi|5868837
                                                                                 4.3
        329362
                                       CH.16_p2 gi|6165201
25
                                                                                 5.5
        329929
                                                                                 7.6
        329960
                                       CH.16_p2 gi|5091594
                                       CH.16_p2 gi|6671887
                                                                                 6
         330020
                                       CH.05_p2 gi|6013592
                                                                                 12.6
        330211
                                       androgen receptor (dihydrotestosterone r
         330384 M23263
                                                                                                                           13.8
30
                                                                                 Antigen, Prostate Specific, Alt. Splice
                                       Hs.321110
         330430 HG2261-HT2352
                             Hs.299867 guanine nucleotide binding protein 4
         330546 U31382
                                                                                 4.9
                                       hepatocyte nuclear factor 3; alpha
         330551 U39840
                             Hs.30732
                                       ESTs
                                                                                 6
         330658 AA319514
                                                                                 5.5
         330700 AA037415
                             Hs.20999
                                       ESTs
                                                                                 5.1
35
                             Hs.6759
                                       ESTs
         330704 AA056557
                             Hs.157078 ESTs
                                                                                 11.7
         330705 AA102571
                             Hs.177576 ESTs; Moderately similar to kynurenine a
                                                                                  14.5
         330706 AA121140
                             Hs.52620
                                       ESTs
         330712 AA167269
                                                                                 7.2
                                       ESTs; Weakly similar to !!!! ALU SUBFAMI
         330725 AA252033
                             Hs.24052
40
         330732 AA281092
                             Hs.35254
                                                                                 4.9
                                       ESTs
                             Hs.15251
                                       Human DNA sequence from clone 437M21 on
         330762 AA449677
                                                                                 18.5
                                                                                 4.3
                             Hs.143187 FK506-binding protein 3 (25kD)
         330763 AA450200
                             Hs.11356
                                       ESTs
                                                                                 5.8
         330772 AA479114
                                                                                  4.6
                                        EST
         330786 D60374
45
                             Hs.91202
                                                                                  15.3
                                       ESTs
         330892 AA149579
                                                                                  10.3
         330949 H01458
                             Hs.142896 ESTs
                                                                                  4.4
         330977 H20826
                             Hs.315181 ESTs
                                                                                 11.8
                             Hs.108920 ESTs
         331017 N24619
                             Hs.14846 ESTs
                                                                                  11.6
         331099 R36671
50
                                                                                  4.8
                             Hs.268714 ESTs
         331128 R51361
                                                                                  13
                             Hs.268838 ESTs
         331151 R82331
                                                                                  4.9
                             Hs.168439 ESTs
         331195 T64447
         331320 AA262999
                             Hs.300141 ESTs
                                                                                  4.8
                                                                                 6.1
                             Hs.87929 ESTs
         331321 AA278355
55
                                                                                  92
                             Hs.118630 ESTs
         331337 AA287662
                             Hs.88143
                                       ESTs
                                                                                  9.9
         331348 AA400596
                                                                                  4.3
                             Hs.81897
                                       ESTs
         331359 AA416979
                                                                                  4.6
         331383 AA454543
                             Hs.43543 ESTs
                             Hs.237339 ESTs; Moderately similar to IIII ALU SUB
                                                                                  4.9
         331422 F10802
                             Hs.41223
                                                                                  7.5
60
                                       ESTs
         331442 H77381
         331466 N21680
                             Hs.43455
                                       ESTs
                                                                                  5.4
                             Hs.44076
                                        ESTs
                                                                                  6.5
         331479 N27154
                             Hs.291039 ESTs; Weakly similar to hypothetical 43.
                                                                                  12.5
         331490 N32912
         331493 N34357
                             Hs.93817 ESTs
                                                                                  4.6
65
                             Hs.48703
                                        ESTs
                                                                                  9.2
         331561 N62780
                                                                                  4.6
         331615 N92352
                             Hs.5472
                                        ESTs
         331659 W48868
                             Hs.334305 ESTs
                                                                                  8.7
                             Hs.65949 KIAA0888 protein
                                                                                  10.3
         331696 Z38907
                                                                                  4.8
         331811 AA404500
                             Hs.187958 ESTs
```

	331848	AA417039	Hs.98268	signal recognition particle 72kD	7.5
		AA429445	Hs.98640	ESTS	6.5
		AA431407	Hs.98802	Homo sapiens Chromosome 16 BAC clone CIT	33.6
5		AA460158	Hs.99589 Hs.105322	KIAA1028 protein	6.8 5.3
5		AA464518 AA490831	Hs.201591	FSTe	10.8
		AA599477	Hs.291156		4.4
	332173		Hs.100725		5.5
- C-2		N58172		ESTs	14.2
10	332249	N62096	Hs.194140		7.2
		T79428	Hs.339667		5.6
		AA340504	LI- 007704	ESTs; Weakly similar to similar to human	21.2 15.3
		N75542 N95495	Hs.56729	transcription factor 4 ESTs; Highly similar to GTP-binding prot	7.1
15		L38503		glutathione S-transferase theta 2	6.6
10		AA281753	Hs.17731	inositol 1;4;5-triphosphate receptor; ty	5.8
		M31682	Hs.19280	inhibin; beta B (activin AB beta polypep	5.5
		M99487		folate hydrolase (prostate-specific memb	38.1
20		N48715	Hs.20991	ESTs	6.5
20		D84454	Hs.22587	solute carrier family 35 (UDP-galactose methyl CpG binding protein 2	4.8 5.6
		AA279313 AA412405	Hs.32951 Hs.40513	ESTs; Weakly similar to BETA GALACTOSIDA	5.6
		N95742	Hs.6390	ESTs	6.9
		T94885	Hs.75725	carboxypeptidase E	24.3
25		D26070	Hs.79306	inositol 1:4:5-triphosphate receptor; ty	9.9
		L00058	Hs.79630	v-myc avian myelocytomatosis viral oncog	5.6
		R72029	Hs.83428	synaptophysin-like protein	5
	332781	AA233258		ESTs; Weakly similar to D1007.5 [C.elega CH22_FGENES.6_2	4.5 30.8
30	332798			CH22_FGENES.6_5	66.8
50	332799			CH22_FGENES.6_6	19.8
	332933			CH22_FGENES.38_7	5.6 .
	332980			CH22_FGENES.54_1	5.5
35	332984 333168			CH22_FGENES.54_6 CH22_FGENES.94_1	4.9 4.7
33	333169			CH22_FGENES.94_2	4.4
	333452			CH22_FGENES.157_1	4.8
	333456	•		CH22_FGENES.157_5	4.3
40	333458			CH22_FGENES.157_7	4.6 4.7
40	333611 333621			CH22_FGENES.217_6 CH22_FGENES.219_5	5.5
	333814			CH22_FGENES.282_2	7.1
	333849			CH22_FGENES.290_8	6.2
	333949			CH22_FGENES.303_5	4.3
45	333951			CH22_FGENES.303_7	4.9
	333955			CH22_FGENES.303_11	5.6 5.1
	334150 334223			CH22_FGENES.339_1 CH22_FGENES.360_4	20.3
	334297			CH22_FGENES.372_3	9.4
50	334443			CH22_FGENES.387_2	4.6
	334444			CH22_FGENES.387_4	5.6
	334447			CH22_FGENES.387_7	13.1
	334570			CH22_FGENES.405_11	5.4
55	334749 334777			CH22_FGENES.427_1 CH22_FGENES.430_9	5.3 4.7
33	334960	•		CH22 FGENES.465_29	5.2
	335179			CH22_FGENES.504_9	8.8
	335293			CH22_FGENES.527_6	4.7
60	335550			CH22_FGENES.576_11	5.1
60	335581 335586			CH22_FGENES.581_19 CH22_FGENES.581_25	5.7 4.3
	335809			CH22_FGENES.617_6	6.2
	335810			CH22_FGENES.617_7	5.8
	335822			CH22_FGENES.619_7	7.1
65	335824			CH22_FGENES.619_11	8.5
	335853 335886			CH22_FGENES.626_5 CH22_FGENES.632_4	4.3 4.3
	336034			CH22_FGENES.678_5	6.8
	336441			CH22_FGENES.827_7	7.6

	336624	CH22_FGENES.6-3	43.3
	336625	CH22_FGENES.6-4	37.9
	336679	CH22_FGENES.43-7	5.3
	337577	CH22_C65E1.GENSCAN.8-1	4.9
5	338255	CH22_EM:AC005500.GENSCAN.276-3	13.4
-	338260	CH22_EM:AC005500.GENSCAN.279-10	4.6
	338561	CH22_EM:AC005500.GENSCAN.421-5	4.6
	338562	CH22 EM:AC005500.GENSCAN.421-6	4.3
	338759	CH22_EM:AC005500.GENSCAN.517-6	5.1
10	338763	CH22_EM:AC005500.GENSCAN.517-16	5.5
	338764	CH22_EM:AC005500.GENSCAN.517-17	7.1

TABLE 3A shows the accession numbers for those primekeys lacking unigeneID's for Table 3. For each probeset we have listed the gene cluster number from which the oligonucleotides were designed. Gene clusters were compiled using sequences derived from Genbank ESTs and mRNAs. These sequences were clustered based on sequence similarity using Clustering and Alignment Tools (DoubleTwist, Oakland California). The Genbank accession numbers for sequences comprising each cluster are listed in the "Accession" column.

Pkey: Unique Eos probeset identifier number
CAT number: Gene duster number
Accession: Genhank accession numbers

5

Accession:		on: Ger	Genbank accession numbers					
15	Pkey	CAT number		Accession				
	123619	371681_1		AA602964 AA609200				
		143512_1		Z24878 AA494098 F13654 AA494040 AA143127				
		41847_1		Z83806 AJ132091 AJ132090				
20		1589048_1		H48372 W01626				
		genbank_AA21362		AA213620	•			
		genbank_W38419		W38419				
		entrez_M21305		M21305				
		genbank_N22401		N22401				
25		genbank_AA13659		AA136590				
		47271_1		W69304 AF086283 W69200	•			
		350959_1		Al821085 AW973464 AA554802 Al821831 AA657438 AA640756 AA650339				
		262792 1		AW177009 Al381610				
		300543 1		AA884766 AW974271 AA592975 AA447312				
30		336411_1		Al685464 AW971336 AA513587 AA525142				
		37699_1		Al685464 AW971336 AA513587 AA525142 AF199613 AF108756 Al692552 Al393343 Al800510 Al377711 F24263 AA661876				
		398093_1		Al692552 Al393343 Al800510 Al377711 F24263 AA661876	•			
		376239_1		AA613792 AW182329 T05304 AW858385				
	329362	c_x_hs		, , , , , , , , , , , , , , , , , , ,				
35		CH22_4071FG_6_	3 .					
		CH22_4072FG_6_						
		CH22_4157FG_43						
		CH22_6856FGL						
		CH22 6863FG_L						
40	329929	c16_p2	_					
	329960							
	338561	CH22_7294FGL	INK_EM:AC00					
		CH22_7295FGL						
	338759	CH22_7581FGL	INK_EM:AC00					
45		CH22_7585FGL						
		CH22_7586FGL						
		CH22_400FG_94_						
		CH22_401FG_94_						
50		CH22_702FG_157						
50		CH22_706FG_157						
•		CH22_708FG_157						
		CH22_872FG_217						
		CH22_882FG_219						
EE .		CH22_1083FG_28						
55 ·		CH22_1118FG_29						
		CH22_2515FG_50						
		CH22_1225FG_303						
		CH22_1227FG_303						
60		CH22_1231FG_303		·				
UU		CH22_2635FG_523	/_O_LINK_EM					
	326816							
	326997		6 14 1 MW F					
		CH22_2905FG_576 CH22_2938FG_58						
65		CH22_2936FG_58						
UJ	333300	OI 155_50441 12_00	ELINI\E					

	328492	c 7 hs	
		CH22_3181FG 617 6_LINK_EM	
		CH22_3182FG_617_7_LINK_EN	
_		CH22_3195FG_619_7_LINK_EN	
5		CH22_3197FG_619_11_LINK_E	
		CH22_3228FG_626_5_LINK_EN	
	335886	CH22_3261FG_632_4_LINK_EN	
	330020	c16_p2	
	330211	c_5_p2	
10	337577	CH22 5864FG LINK_C65E1.G	
	307848	Al364186	
		CH22 13FG 6 2 LINK C4G1.G	
		CH22_14FG_6 5_LINK_C4G1.G	
		CH22 15FG 6 6 LINK C4G1.G	
15		CH22 1429FG 339 1 LINK EM	
IJ		CH22_154FG_38_7_LINK_C20H	
		CH22_204FG_54_1_LINK_EM:A	
			·
		CH22_208FG_54_6_LINK_EM:A	
20		CH22_1507FG_360_4_LINK_EN	
20		CH22_1588FG_372_3_LINK_EN	· ·
		c21_hs	
		CH22_1742FG_387_2_LINK_EN	
		CH22_1743FG_387_4_LINK_EN	
~~		CH22_1746FG_387_7_LINK_EN	
25		CH22_1875FG_405_11_LINK_E	
	334749	CH22_2061FG_427_1_LINK_EM	1
	334777	CH22_2089FG_430_9_LINK_EN	1
		CH22_3419FG_678_5_LINK_DJ	
	334960	CH22_2281FG_465_29_LINK_E	
30	336441	CH22_3861FG_827_7_LINK_DJ	
	330551	9851_2	U39840 NM_004496 AW135607 BE087458 BE087567 AA177116 AW195705 AW750756 AI811008 AI694151
			BE348594 AW971075 Al347950 Al201455 Al073898 AA652680 AA613671 Al318364 AA507550 AA693692
			Al032599 AA991871 Al269801 AW948974 T74639 AA532907 AW949173
	330786	53973_3	BE379594 Al192455 AL039862 Al744012 Al761735 AW243181 Al743687 Al928223 Al423022 Al627855
35		- · · · · · -	AI636059 AI651571 AW802044 AI826995 AI431733 AI539125 AA863056 AW270910 AI768930 AW008835
			AW615183 AW591147 Al695294 Al672106 AA506358 Al308060 AA011556 AA962437 Al935488 BE219625
			AI004356 AW151394 AI218466 N66178 AI419784 AW242519 AW946907 D60374 AA989263 AI698799
		•	AA470460 Al824167
	332247	372969_1	AA669097 AA513815 AA026798 AA676526 AA704429 AA704269 AW118292 AA579216 N58172
40		20265_1	AW579842 BE156562 BE156690 BE156489 BE081033 AK001559 BE149402 M85387 AW367811 AW367798
	0000	2020-	R17370 Al908947 AA382932 R58449 H18732 AA371231 AW962899 AA713530 AW892946 R53463 H11063
			AW068542 Z40761 BE176212 BE176155 W23952 W92188 AW374883 AA303497 AW954769 AA036808
			BE168063 AW382073 AW382085 AL041475 H80748 Al078161 BE463983 Al805213 Al761264 W94885
			N94502 Al623772 Al419532 Al610302 Al634190 AW002516 AW150777 Al352312 Al367474 AW204807
45			AI675502 AI337026 AW134715 BE328451 AI123157 AI560020 AI300745 AI608631 AI248873 AA742484
40			AW051635 H18646 Al245045 AA507111 Al640510 Al925594 AA115747 AA143035 AA151106
	000704	20044-4	AK001764 BE313896 AA380199 AA380151 AA194996 AW118089 AA495871 AW975219 AW085598
	332/81	32044_1	MINOR DESCRIPTION AND DESCRIPTION AND DESCRIPTION AND ADDRESS AND
			AI378909 AW992310 AW992409 AI911857 AA657643 AI804471 AI242589 AI623968 R09556 AI129100
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50			AA001729 W90790 BE090656 AW295015 Al674596 Al431734 Al420517 AW769185 Al128355 Al192474
			AI820001 AA001929 AA706925 AI076676 AI499119 AI200493 AI695919 AI376217 W69195 W69261
			AW305099 W90320 BE048357 Al658856 AA838534 AA233258 Al753393 AA709227 Al674387 Al872616
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TABLE 3B shows the genomic positioning for those primekeys lacking unigene ID's and accession numbers in Table 3. For each predicted exon, we have listed the genomic sequence source used for prediction. Nucleotide locations of each predicted exon are also listed.

10		Unique number corresponding to an Eos probeset Sequence source. The 7 digit numbers in this column are Genbank Identifier (GI) numbers. "Dunham I. et al." refers to the DNA sequence of human chromosome 22." Dunham I. et al., Nature (1999) 402:489-495. Indicates DNA strand from which exons were predicted. Indicates nucleotide positions of predicted exons.)
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332980 Dunham, I. et.al. Minus 5136165-5136019	
333168 Dunham, I. et.al. Minus 3729896-3729788	
333169 Dunham, I. et.al. Minus 3730864-3730767	
0 333452 Dunham, I. et.al. Minus 5136165-5136019	
333456 Dunham, I. et.al. Minus 2631933-2631797	
333458 Dunham, I. et.al. Minus 5143942-5143806	
334223 Dunham, I. et.al. Minus 12734365-12734269	
334749 Dunham, I. et.al. Minus 16090686-16090106	
335293 Dunham, I. et.al. Minus 22316408-22316275	
335550 Dunham, I. et.al. Minus 24668714-24668658	
335853 Dunham, I. et.al. Minus 26614629-26614506	
336624 Dunham, I. et.al. Minus 227714-227577	
336625 Dunham, I. et.al. Minus 229124-229024	
336679 Dunham, I. et.al. Minus 2035790-2035681	
338255 Dunham, I. et.al. Minus 15242294-15242231	
338561 Dunham, I. et.al. Minus 22311966-22311856	
338562 Dunham, I. et.al. Minus 22312594-22312465	
5 338759 Dunham, I. et.al. Minus 26582475-26582199	
338763 Dunham, I. et.al. Minus 26628148-26628009 338764 Dunham, I. et.al. Minus 26641232-26641101	

	329900	5091594	Minus	1031-1102
	329929	6165201	Minus	156410-156553
	330020	6671887	Plus	172397-172491
	326816	6552458	Plus	198354-198436
5	326997	5867660	Minus	71389-72147
	327098	6682516	Minus	1061684-1062361
	330211	6013592	Plus	59158-59215
	328492	5868455	Minus	46094-46241
	329362	5868837	Minus	65688-68173
10				

TABLE 4: shows a preferred subset of the Accession numbers for genes found in Table 3 which are differentially expressed in prostate tumor tissue compared to normal prostate tissue.

Pkey: ExAcon: Unique Eos probeset identifier number Exemplar Accession number, Genbank accession number UnigenelD: Unigene Title: R1: Unigene number Unigene gene title

Ratio of tumor to normal body tissue

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15	Pkey	ExAccn	UnigenelD	Unigene Title	R1
	100819	HG4020-HT42	90Hs.2387	Transglutaminase	10.5
	102698	U75272	Hs.1867	progastricsin (pepsinogen C)	10.6
	102869	X02544	Hs.572	orosomucoid 1	22.6
	105370	AA236476	Hs.22791	ESTs; Weakly similar to transmembrane pr	10.3
20	105645	AA282138	Hs.11325		14
	106094	AA419461	Hs.23317	ESTs	10.9
	109014	AA156790	Hs.262036	ESTs	15.3
	109562	F01811	Hs.187931	ESTs; Moderately similar to voltage-gate	10.8
	113021	T23855		KIAA1028 protein	10.8
25	114124	Z38595	Hs.125019	ESTs; Highly similar to KIAA0886 protein	21.3
	122791	AA460158		KIAA1028 protein	12.4
	124352	N21626	Hs.102406	ESTs	10.2
	301042	Al659131	Hs.197733	ESTs	24.9
	302005	A1869666	Hs.123119		36.8
30 .	302410	NM_004917	Hs.218366	EST cluster (not in UniGene) with exon h	26.8
	302881	AA508353		relaxin 1 (H1)	78.8
	303344	AA255977		ESTs; Highly similar to ubiquitin-conjug	19.5
	303753	AW503733	Hs.9414	ESTs	13
	310431	Al420227	Hs.149358		72.9
35	311251	AI655662	Hs.197698	ESTs	41.3
	311596	A1682088	Hs.79375	ESTs	26.4
		AA759250		cytochrome b-561	11
		AA033609	Hs.239884		11.2
		AA861697		EST duster (not in UniGene)	13.4
40		Al821895	Hs.193481		29.4
		Al672225	Hs.222886		19.3
		AW292425	Hs.163484		15.5
		AA876910	Hs.134427		20
45		AI654187	Hs.195704		14.2
45		AW295184		ESTs; Weakly similar to DEOXYRIBONUCLE	
		AI949409	Hs.194591		12.3
		AW291511	Hs.159066		25.9
	319080			ESTs	16.9
50		AA460775	Hs.6295	ESTs	14.3
50		AF071202		ATP-binding cassette; sub-family C (CFTR	56.2
		AW297633	Hs.118498		14.7
		W07459		EST duster (not in UniGene)	22
		AA056060		EST duster (not in UniGene)	18.4
55		AW043782	Hs.293616		10.7
22		AA639902	Hs.104215		24.7
		AW016378 AA508552	Hs.292934 Hs.195839		24.2 54
		AASU8552 AI694767	Hs.129179		22
		Al217963		ESTs; Weakly similar to Pro-a2(XI) [H.sa	10.6
60		AA641092	Hs.257339		10.2
OU		AI557019	Hs.116467		34.4
	330211	MI33/018	115.110407	CH.05_p2 gi 6013592	12.6
		Degree uma	50 Un 20111	O Antigen, Prostate Specific, Alt. Splice	13.8
		AA121140	UC 177574	ESTs; Moderately similar to kynurenine a	14.5
65		AA449677			
UJ		AA149579	Hs.91202		15.3
		H01458	Hs.142896		10.3
	330348	1101400	113.142.090	LUIS	1023

	331099	R36671	Hs.14846	ESTs	11.6
	331151	R82331	Hs.268838	ESTs	13
	331889	AA431407	Hs.98802	Homo sapiens Chromosome 16 BAC clone	CIT 33.6
	332247	N58172		ESTs	14.2
5	332396	AA340504		ESTs; Weakly similar to similar to human	21.2
	332533	M99487	Hs.325825	folate hydrolase (prostate-specific memb	38.1
	332697	T94885	Hs.75725	carboxypeptidase E	24.3
	332797			CH22_FGENES.6_2	30.8
	332798			CH22_FGENES.6_5	66.8
10	332799			CH22_FGENES.6_6	19.8
	334223			CH22_FGENES.360_4	20.3
	336624			CH22_FGENES.6-3	43.3
	336625			CH22 FGENES.6-4	37.9

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TABLE 4A shows the accession numbers for those primekeys lacking unigeneID's for Table 4. For each probeset we have listed the gene cluster number from which the oligonucleotides were designed. Gene clusters were compiled using sequences derived from Genbank ESTs and mRNAs. These sequences were clustered based on sequence similarity using Clustering and Alignment Tools (DoubleTwist, Oakland California). The Genbank accession numbers for sequences comprising each cluster are listed in the "Accession" column.

10	Pkey: Unique Eos probeset CAT number: Gene cluster number Accession: Genbank accession n				
15	Pkey CAT	number	Accession		
20	336625 CH2: 330211 c_5_ 332797 CH2: 332798 CH2: 332799 CH2:	2_4071FG_6_3_ 2_4072FG_6_4_ p2 2_13FG_6_2_LINK_C4G1.G 2_14FG_6_5_LINK_C4G1.G 2_15FG_6_6_LINK_C4G1.G 2_1507FG_360_4_LINK_EM	· · · · · · · · · · · · · · · · · · ·		
25	332247 3729 332396 2026	69_1	AA669097 AA513815 AA026798 AA676526 AA704429 AA704269 AW118292 AA579216 N58172 AW579842 BE156562 BE156690 BE156489 BE081033 AK001559 BE149402 M85387 AW367811 AW367798 R17370 Al908947 AA382932 R56449 H18732 AA371231 AW962899 AA713530 AW892946 R53463 H11063 AW068542 Z40761 BE176212 BE176155 W23952 W92188 AW374883 AA303497		
30			AW954769 AA036808 BE168063 AW382073 AW382085 AL041475 H80748 AI078161 BE463983 Al805213 AI761264 W94885 N94502 AI623772 AI419532 AI810302 AI634190 AW002516 AW150777 AI352312 AI367474 AW204807 AI675502 AI337026 AW134715 BE328451 AI123157 AI560020 AI300745 AI608631 AI248873 AA742484 AW051635 H18646 AI245045 AA507111 AI640510 AI925594 AA115747 AA143035 AA151106		

TABLE 4B shows the genomic positioning for those primekeys lacking unigene ID's and accession numbers in Table 4. For each predicted exon, we have listed the genomic sequence source used for prediction. Nucleotide locations of each predicted exon are also listed.

10	Pkey: Ref:		Sequence source. The DNA sequence of hum	Unique number corresponding to an Eos probeset Sequence source. The 7 digit numbers in this column are Genbank Identifier (GI) numbers. "Dunham I. et al." refers to the publication entitled "The DNA sequence of human chromosome 22." Dunham I. et al., Nature (1999) 402:489-495.				
	Strand: Nt_position:			d from which exons were predicted. positions of predicted exons.				
15	Pkey	Ref	Strand	Nt_position				

15	Pkey	Ref	Strand	Nt_position	
20	332797 332798 332799 334223 336624 336625 330211	Dunham, I. et.al. Dunham, I. et.al. Dunham, I. et.al. Dunham, I. et.al. Dunham, I. et.al. Dunham, I. et.al. 6013592	Minus Minus Minus Minus Minus Minus Plus	216964-216798 232147-231974 232421-232307 12734365-12734269 227714-227577 229124-229024 59158-59215	

TABLE 5: 1170 GENES UP-REGULATED IN PROSTATE CANCER COMPARED TO NORMAL ADULT TISSUES

Table 5 shows 1170 genes up-regulated in prostate cancer compared to normal adult tissues. These were selected from 59680 probesets on the Affymetrix/Eos Hu03 GeneChip array such that the ratio of "average" prostate cancer to "average" normal adult tissues was greater than or equal to 3.44. The "average" prostate cancer level was set to the 85th percentile amongst 73 prostate cancers. The "average" normal adult tissue level was set to the 85th percentile amongst 162 non-malignant tissues. In order to remove gene-specific background levels of non-specific hybridization, the 7.5th percentile value amongst the 162 non-malignant tissues was subtracted from both the numerator and the denominator before the ratio was evaluated.

15	Pkey: ExAccn: UnigeneID:	Unique Eos probeset identifier number Exemplar Accession number, Genbank accession number Unigene number Unigene gene title
	Unigene Title: R1:	Ratio of tumor to normal tissue

20	Pkey	ExAccn	UnigeneiD	Unigene Title .	R1
	446057	AJ420227	Hs.149358	ESTs. Weakly similar to A46010 X-linked	86.42
	400302	N48056	Hs.1915	folate hydrolase (prostate-specific memb	66.46
	414569	AF109298	Hs.118258	prostate cancer associated protein 1	58.36
25	417407	AA923278	Hs.290905	ESTs, Weakly similar to protease [H.sapi	56.16
	431579	AW971082	Hs.222886	ESTs, Weakly similar to TRHY_HUMAN TRICH	53.38
	409361	NM_005982	Hs.54416	sine oculis homeobox (Drosophila) homolo	48.28
•	409731	AA125985	Hs.56145	thymosin, beta, identified in neuroblast	45.24
	400298	AA032279	Hs.61635	six transmembrane epithelial antigen of	43.48
30	420154	Al093155	Hs.95420	JM27 protein	41.12
	433466	AA508353	Hs.105314	retaxin 1 (H1)	39.88
	400296	AA305627	Hs.139336	ATP-binding cassette, sub-family C (CFTR	38.42
	400292	AA250737	Hs.72472	ESTs	38.00
	432887	Al926047	Hs.162859	ESTs	36.48
35	439176	A1446444	Hs.190394	ESTs, Weakly similar to B28096 line-1 pr	36.45
	430722	AW968543	Hs.203270	ESTs, Weakly similar to ALU1_HUMAN ALU S	33.20
	437052	AA861697	Hs.120591	ESTs	33.02
	418396	A1765805	Hs.26691	ESTs	32.68
	434036	Al659131	Hs.197733	hypothetical protein MGC2849	32.44
40	407709	AA456135	Hs.23023	ESTs	32.10
	426747	AA535210	Hs.171995	kallikrein 3, (prostate specific antigen	31.80
	407168	R45175		ESTs	31.72
	440260	Al972867	Hs.7130	copine IV	30.52
	421513	X00949	Hs.105314	relaxin 1 (H1)	30.10
45	416370	N90470	Hs.203697	ESTs, Weakly similar to 138022 hypotheti	29.68
	407122	H20276	Hs.31742	ESTs -	29.24
	400287	S39329	Hs.181350	kallikrein 2, prostatic	28.90
	432244	A1669973	Hs.200574	ESTs	28.74
~^	451939	U80456	Hs.27311	single-minded (Drosophila) homolog 2	28.74
50	415989	Al267700	Hs.111128	ESTs	28.34
	418961	AW967646	Hs.23023	ESTs	27.34
	425628	NM_004476	Hs.1915	folate hydrolase (prostate-specific memb	27.32
	458509	AA654650	Hs.282906	ESTs	27.24
	448290	AK002107	Hs.20843	Homo sapiens cDNA FLJ11245 fis, clone PL	27.16
55	428336	AA503115	Hs.183752	microseminoprotein, beta-	26.17
	450096	Al682088	Hs.223368	holocarboxylase synthetase (biotin-[prop	25.60
	400299	X07730	Hs.171995	kallikrein 3, (prostate specific antigen	24.91
	437571	AA760894	Hs.153023	ESTs	24.74
60	453160	Al263307	Hs.146228	H2B histone family, member L	24.66
60	453096	AW294631	Hs.11325	ESTs	24.46
	425075	AA506324	Hs.1852	acid phosphatase, prostate	24.23
	407202	N58172	Hs.109370	ESTs	24.18

	424846	AU077324	Hs.1832	neuropeptide Y	23.57
	453370	A1470523	Hs.182356	ATP-binding cassette, sub-family C (CFTR	23.16
	422805	AA436989	Hs.121017	H2A histone family, member A	22.52
5	444917	R68651	Hs.144997	ESTs	22.26
3	408826	AF216077	Hs.48376	Homo sapiens clone HB-2 mRNA sequence	22.02
	413597 426429	AW302885	Hs.117183	ESTS	21.76 21.32
	435981	X73114 H74319	Hs.169849 Hs.188620	myosin-binding protein C, slow-type ESTs	21.12
	432966	AA650114	115.100020	ESTs	21.12
10	418848	Al820961	Hs.193465	ESTs	21.06
	405685	, ,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	1.000		20.90
	443271	BE568568	Hs.195704	ESTs	19.98
	418819	AA228776	Hs.191721	ESTs	19.94
	420757	X78592	Hs.99915	androgen receptor (dihydrotestosterone r	19.72
15	418994	AA296520	Hs.89546	selectin E (endothelial adhesion molecul	19.56
	429918	AW873986	Hs.119383	ESTs	19.04
	415539	Al733881	Hs.72472	ESTs	18.43
	450382	AA397658	Hs.60257	Homo sapiens cDNA FLJ13598 fis, clone PL	18.34
20	418829 429984	AA516531 AL050102	Hs.55999 Hs.227209	NK homeobox (Drosophila), family 3, A hypothetical protein FLJ21617	18.28 17.82
20	443822	A1087412	Hs.143611	ESTs, Weakly similar to 2004399A chromos	17.66
	431676	Al685464	Hs.292638	gb:tt88f04.x1 NCI_CGAP_Pr28 Homo sapiens	17.64
	410330	AW023630	Hs.46786	ESTs	17.52
	432441	AW292425	Hs.163484	ESTs	17.A1
25	452792	AB037765	Hs.30652	KIAA1344 protein	17.39
	445472	AB006631	Hs.12784	Homo sapiens mRNA for KIAA0293 gene, par	17.00
	414565	AA502972	Hs.183390	hypothetical protein FLJ13590	16.82
	430487	D87742	Hs.241552	KIAA0268 protein	16.72
30	431716 419536	D89053 AA603305	Hs.268012	fatty-acid-Coenzyme A ligase, long-chain gb:np12d11.s1 NCI_CGAP_Pr3 Homo sapiens	16.60 16.50
50	439677	R82331	Hs.164599	ESTs	16.46
	449625	NM_014253	Hs.23796	odz (odd Oz/ten-m, Drosophila) homolog 1	16.32
	408430	S79876	Hs.44926	dipeptidylpeptidase IV (CD26, adenosine	16.28
	447033	Al357412	Hs.157601	ESTs	16.02
35	453006	Al362575	Hs.167133	ESTs	15.74
	431474	AL133990	Hs.190642	ESTs	15.70
	420218	AW958037	Hs.22437	ribosomal protein L4	15.64
	408000 416208	L11690 AW291168	Hs.620 Hs.41295	bullous pemphigoid antigen 1 (230/240kD) ESTs, Weakly similar to MUC2_HUMAN MUCIN	15.54 15.48
40	430226	BE245562	Hs.2551	adrenergic, beta-2-, receptor, surface	15.40
•••	415263	AA948033	Hs.130853	ESTs	15.38
	432437	W07088	Hs.293685	ESTs	15.26
	428398	Al249368	Hs.98558	ESTs	15.21
45	429900	AA460421	Hs.30875	ESTs	14.90
45	449156	AF103907	Hs.171353	prostate cancer antigen 3	14.89
	411096 435974	U80034	Hs.68583	mitochondrial intermediate peptidase	14.81
	444484	U29690 AK002126	Hs.37744 Hs.11260	Homo sapiens beta-1 adrenergic receptor hypothetical protein FLJ11264	14.76 14.76
	422728	AW937826	Hs.103262	ESTs, Weakly similar to ZN91_HUMAN ZINC	14.60
50	418601	AA279490	Hs.86368	calmegin	14.56
	448999	AF179274	Hs.22791	transmembrane protein with EGF-like and	14.55
	445885	AI734009	Hs.127699	KIAA1603 protein	14.44
	452712	AW838616		gb:RC5-LT0054-140200-013-D01 LT0054 Homo*	14.22
55	432189	AA527941	(1- ======	gb:nh30c04.s1 NCI_CGAP_Pr3 Homo saplens	14.12
23	424565	AW102723	Hs.75295	guanylate cyclase 1, soluble, alpha 3 neurofilament, heavy polypeptide (200kD)	13.78
	429290 419264	AF203032 AA877104	Hs.198760 Hs.293672	ESTs, Weakly similar to ALUB_HUMAN !!!!	13.57 13.40
	416445	AL043004	Hs.300678	KIAA0135 protein	13.40
	407275	Al364186	110.000070	gb:gw34h07.x1 NCI_CGAP_Ut4 Homo sapiens	13.24
60	408369	R38438	Hs.182575	solute carrier family 15 (H+/peptide tra	13.21
	446720	Al439136	Hs.140546	ESTs	13.06
	434988	Al418055	Hs.161160	ESTs	13.02
	448172	N75276	Hs.135904	ESTs	12.98
65	416182	NM_004354	Hs.79069	cyclin G2	12.94
UJ	420544	AA677577	Hs.98732	Homo saplens Chromosome 16 BAC clone CIT	12.79
	445413 452588	AA151342 AA889120	Hs.12677 Hs.110837	CGI-147 protein homeo box A10	12.64
	407819	R42185	Hs.274803	ESTs	12.62 12.60
	433444	AW975324	Hs.129816	ESTs	12.60

	421059	Al654133	Hs.30212	thyroid receptor interacting protein 15	12.30
	420077	AW512260	Hs.87767	ESTs	12.24
	453930	AA419466	Hs.36727	hypothetical protein FLJ10903	12.22
	441610				12.20
_		AW576148	Hs.148376	ESTs	
- 5	451009	AA013140	Hs.115707	ESTs	12.18
	433764	AW753676	Hs.39982	ESTs	12.16
	440286	U29589	Hs.7138	cholinergic receptor, muscarinic 3	12.04
	443912	R37257	Hs.184780	ESTs	11.92
		Al821895	Hs.193481	ESTs	11.91
10	419526			=	
10	423073	BE252922	Hs.123119	MAD (mothers against decapentaplegic, Dr	11.87
	452784	BE463857	Hs.151258	hypothetical protein FLJ21062	11.86
	414422	AA147224	Hs.71814	ESTs	11.76
	450203	AF097994	Hs.301528	L-kynurenine/alpha-aminoadipate aminotra	11.68
	436679	Al127483	Hs.120451	ESTs, Weakly similar to unnamed protein	11.60
15					11.60
13	440901	AA909358	Hs.128612	ESTs	
	448045	AJ297436	Hs.20166	prostate stem cell antigen	11.51
	433887	AW204232	Hs.279522	ESTs	11.50
	434980	AW770553	Hs.293640	sterol O-acyltransferase (acyl-Coenzyme	11.38
	425905	AB032959	Hs.161700	novel C3HC4 type Zinc finger (ring finge	11.33
20				ESTs	11.32
20	434680	T11738	Hs.127574		
	449650	AF055575	Hs.297647	calcium channel, voltage-dependent, L ty	11.18
	431173	AW971198	Hs.294068	ESTs	11.16
	434539	AW748078	Hs.214410	ESTs, Weakly similar to MUC2_HUMAN MUCIN	11.16
	410037	AB020725	Hs.58009	KIAA0918 protein	11.14
25			Hs.50495	ESTs	11.14
23	417708	N74392			
	458332	Al000341	Hs.220491	ESTs	11.12
	420381	D50640	Hs.301782	phosphodiesterase 3B, cGMP-inhibited	11.10
	425665	AK001050	Hs.159066	hypothetical protein FLJ10188	11.08
	425710	AF030880	Hs.159275	solute carrier family, member 4	11.08
30			Hs.191381	hypothetical protein	11.04
20	428728	NM_016625	H2.131301		
	407021	U52077		gb:Human mariner1 transposase gene, comp	11.02
	410733	D84284	Hs.66052	CD38 antigen (p45)	11.02
	401714				10.90
	434485	Al623511	Hs.118567	ESTs	10.89
35	415786	AW419196	Hs.257924	hypothetical protein FLJ13782	10.87
33					10.85
	452340	NM_002202	Hs.505	ISL1 transcription factor, LIM/homeodoma	
	453628	AW243307	Hs.170187	hypothetical protein	10.72
	408063	BE086548	Hs.42346	calcineurin-binding protein calsarcin-1	10.67
	417687	Al828596	Hs.250691	ESTs	10.64
40	434666	AF151103	Hs.112259	T cell receptor gamma locus	10.53
-10			Hs.301885	Homo sapiens cDNA FLJ11346 fis, clone PL	10.50
	432374	W68815			
	428819	AL135623	Hs.193914	KIAA0575 gene product	10.48
	413409	Al638418	Hs.21745	DEAD/H (Asp-Glu-Ala-Asp/His) box polypep .	10.44
	428775	AA434579	Hs.143691	ESTs	10.21
45	436556	Al364997	Hs.7572	ESTs	10.20
	441690	R81733	Hs.33106	ESTs	10,14
				hypothetical protein dJ551D2.5	10.10
	419852	AW503756	Hs.286184		
	421991	NM_014918	Hs.110488	KIAA0990 protein	10.04
	423698	AA329796	Hs.1098	DKFZp434J1813 protein	10.02
50	452039	A1922988	Hs.172510	ESTs	10.00
_	433043	W57554	Hs.125019	ESTs	9.98
	433927	Al557019	Hs.116467	small nuclear protein PRAC	9.97
				and the CHO demain hinding metals	9.96
	445424	AB028945	Hs.12696	cortactin SH3 domain-binding protein	
	432240	A1694767	Hs.129179	Homo sapiens cDNA FLJ13581 fis, clone PL	9.88
55	433104	AL043002	Hs.128246	ESTs, Moderately similar to unnamed prot	9.84
	452744	A1267652	Hs.30504	Homo sapiens mRNA; cDNA DKFZp434E082 (fr	9.82
	431217	NM_013427	Hs.250830	Rho GTPase activating protein 6	9.75
		_		chromosome 21 open reading frame 11	9.70
	427398	AW390020	Hs.20415		
C C	446896	T15767	Hs.22452	Homo saplens mRNA for KIAA1737 protein,	9.70
60	421470	R27496	Hs.1378	annexin A3	9.64
	406554				9.60
	401424				9.58
		AI 117474	Uc #1101	Homo sapiens mRNA; cDNA DKFZp727C191 (fr	9.56
	407902	AL117474	Hs.41181		
	423545	AP000692	Hs.129781	chromosome 21 open reading frame 5	9.54
65	439024	R96696	Hs.35598	ESTs	9.51
	431548	AI834273	Hs.9711	novel protein	9.48
	409262	AK000631	Hs.52256	hypothetical protein FLJ20624	9.45
					9.42
	446271	D82484	Hs.100469	ESTS	
	448692	AW013907	Hs.224276	methylcrotonoyl-Coenzyme A carboxylase 2	9.26

	414140	AA281279	Hs.23317	hypothetical protein FLJ14681	9.24
	435980	AF274571	Hs.129142	deoxyribonuclease li beta	9.24
	421246	AW582962	Hs.300961	CGI-47 protein	9.20
	427304	AA761526	Hs.163853	ESTs	9.16
5	442914	AW188551	Hs.99519	hypothetical protein FLJ14007	9.16
-	413627	BE182082	Hs.246973	ESTs	9.14
	439699	AF086534	Hs.187561	ESTs, Moderately similar to ALU1_HUMAN A	9.10
	437718	Al927288	Hs.196779	ESTs	9.07
	439820	AL360204	Hs.283853	Homo sapiens mRNA full length insert cDN	9.06
10			Hs.19322	Homo saplens, Similar to RIKEN cDNA 2010	9.05
10	447342	Al199268			9.04
	446223	BE300091	Hs.119699	hypothetical protein FLJ12969	9.03
	410001	AB041036	Hs.57771	kallikrein 11	
	424012	AW368377	Hs.137569	tumor protein 63 kDa with strong homolog	9.03
1.5	441791	AW372449	Hs.175982	hypothetical protein FLJ21159	9.02
15	448206	BE622585	Hs.3731	ESTs, Moderately similar to 138022 hypot	9.02
	414269	AA298489		olfactory receptor, family 51, subfamily	8.99
	442081	AA401863	Hs.22380	ESTs	8.98
	420092	AAB14043	Hs.88045	ESTs	8.85
	411630	U42349	Hs.71119	Putative prostate cancer tumor suppresso	8.80
20	421863	Al952677	Hs.108972	Homo sapiens mRNA; cDNA DKFZp434P228 (fr	8.80
	454141	AW138413	Hs.182356	ATP-binding cassette, sub-family C (CFTR	8.80
	418278	A1088489	Hs.83937	hypothetical protein	8.78
	428330	L22524	Hs.2256	matrix metalloproteinase 7 (matrilysin,	8.76
	432415	T16971	Hs.289014	ESTs, Weakly similar to A43932 mucin 2 p	8.75
25	424906	AI566086	Hs.153716	Homo sapiens mRNA for Hmob33 protein, 3'	8.74
	415245	N59650	Hs.27252	ESTs	8.72
	442409	BE208843	Hs.129544	hypothetical protein MGC15438	8.70
	404571		110.120044	hypotholical protest mad to too	8.66
	418033	W68180	Hs.259855	elongation factor-2 kinase	8.64
30		AW967956	Hs.123648	ESTs, Weakly similar to AF108460 1 ubinu	8.56
50	456497	M11807300	113.120040	LOTS, Weardy Summar to AP 100400 T donid	8.54
	405876	A1574040	Hs.7549	ESTs	8.52
	448807	Al571940			8.48
	445372	N36417	Hs.144928	ESTs	
35	425171	AW732240	Hs.300615	ESTs	8.44
33	419968	X04430	Hs.93913	interleukin 6 (interferon, beta 2)	8.36
	407385	AA610150	Hs.272072	ESTs, Weakly similar to 138022 hypotheti	8.31
	433172	AB037841	Hs.102652	hypothetical protein ASH1	8.30
	422631	BE218919	Hs.118793	hypothetical protein FLJ10688	8.27
40	412719	AW016610	Hs.129911	ESTs	8.24
40	418849	AW474547	Hs.53565	Homo sapiens PIG-M mRNA for mannosyltran	8.22
	444922	Al921750	Hs.144871	Homo sapiens cDNA FLJ13752 fis, clone PL	8.22
	427674	NM_003528	Hs.2178	H2B histone family, member Q	8.20
	432101	Al918950	Hs.11092	EphA3	8.17
	416288	H51299		gb:yp07c06.s1 Soares breast 3NbHBst Homo	8.15
45	404915				8.08
	440106	AA864968	Hs.127699	KIAA1603 protein	8.07
	442861	AA243837	Hs.57787	ESTs	8.06
	452259	AA317439	Hs.28707	signal sequence receptor, gamma (translo	8.06
	443250	AI041530	Hs.132107	ESTs	8.06
50	437267	AW511443	Hs.258110	ESTs	8.04
	452891	N75582	Hs.212875	ESTs, Weakly similar to DYH9_HUMAN CILI	8.02
	422219	AW978073		regulator of mitotic spindle assembly 1	8.00
	453049	BE537217	Hs.30343	EŠTs	8.00
	439731	Al953135	Hs.45140	hypothetical protein FLJ14084	7.98
55	408554	AA836381	Hs.7323	nuclear receptor co-repressor/HDAC3 comp	7.94
	421154	AA284333	Hs.287631	Homo sapiens cDNA FLJ14269 fis, clone PL	7.94
	430107	AA465293	Hs.105069	ESTs	7.94
	433404	T32982	Hs.102720	ESTs	7.93
	450813	Al739625	Hs.203376	ESTs	7.90
60	416239	AL038450	Hs.48948	ESTs	7.85
50			10.40540	gb:tc87d07_x1 NCI_CGAP_CLL1 Homo sapiens	7.82
	448212	AI475858	Un 074500		
	449532	W74653	Hs.271593	ESTs, Moderately similar to A47582 B-cel	7.82
	413930	M86153	Hs.75618	RAB11A, member RAS oncogene family	7.80
65	458191	Al420611	Hs.127832	ESTS	7.80
65	444858	Al199738	Hs.208275	ESTs, Weakly similar to ALUA_HUMAN !!!!	7.78
	457498	Al732230	Hs.191737	ESTs	7.78
	407235	D20569	Hs.169407	SAC2 (suppressor of actin mutations 2, y	7.76
	433759	AA680003	Hs.109363	Homo sapiens cDNA: FLJ23603 fis, done L	7.74
	433805	AA706910	Hs.112742	ESTs	7.74

	400405	AULA COCCOOT	Ha 1700/0	platelet demand growth forter recenter	7.72
	426485	NM_006207	Hs.170040	platelet-derived growth factor receptor- Homo sapiens cDNA FLJ13136 fis, clone NT	7.72
	446028	R44714	Hs.106795		7.70
	418555	Al417215	Hs.87159	hypothetical protein FLJ12577	7.70
_	447499	AW262580	Hs.147674	protocadherin beta 16	7.68
5	419839	U24577	Hs.93304	phospholipase A2, group VII (platelet-ac	7.68
	416857	AA188775	Hs.292453	ESTs	7.66
	413801	M62246	Hs.35406	ESTs, Highly similar to unnamed protein	7.66
	425480	AB023198	Hs.158135	KIAA0981 protein	7.64
10	420120	AL049610	Hs.95243	transcription elongation factor A (SII)-	
10	424099	AF071202	Hs.139336	ATP-binding cassette, sub-family C (CFTR	7.64
	446307	T50083	Hs.9094	ESTs	7.63 7.59
	429220	AW207206	Hs.136319	ESTs	7.54
	420345	AW295230	Hs.25231	ESTS	7.54
15	429208	AA447990	Hs.190478	ESTs Homo sapiens cDNA FLJ13090 fis, clone NT	7.53
15	447247	AW369351	Hs.287955		7.53
	440995	T57773	Hs.10263	ESTs	7.52
	448706	AW291095	Hs.21814	interleukin 20 receptor, alpha exostoses (multiple)-like 2	7.49
	410227	AB009284	Hs.61152	ESTs, Weakly similar to I38022 hypotheti	7.46
20	431616	AA508552	Hs.195839	ESTs. Weakly Sillillai to 130022 hypotheti	7.44
20	434217	AW014795	Hs.23349	Homo sapiens mRNA; cDNA DKFZp434E0528 (f	7.42
	431467	N71831	Hs.256398 Hs.244334	Homo sapiens prostein mRNA, complete cds	7.42
	448519	AW175665	Hs.34981	ESTs	7.40
	446791	A1632278 AW408762	Hs.127478	Homo sapiens clone 24416 mRNA sequence	7.39
25	419743	BE247129	Hs.145569	ESTs	7.36
23	445855 425211	M18667	Hs.1867	progastricsin (pepsinogen C)	7.35
	419131	AA406293	Hs.301622	ESTs	7.34
	400294	N95796	Hs.179809	Homo sapiens prostein mRNA, complete cds	7.33
	441736	AW292779	Hs.169799	ESTs	7.28
30	427701	AA411101	Hs.221750	nuclear autoantigenic sperm protein (his	7.24
50	457733	AW974812	Hs.291971	ESTs	7.24
	418432	M14156	Hs.85112	insulin-like growth factor 1 (somatomedi	7.22
	441201	AW118822	Hs.128757	ESTs	7.21
	419953	BE267154	Hs.125752	ESTs	7.20
35	419991	AJ000098	Hs.94210	eyes absent (Drosophila) homolog 1	7.20
	425018	BE245277	Hs.154196	E4F transcription factor 1	7.20
	424560	AA158727	Hs.150555	protein predicted by clone 23733	7.18
	435380	AA679001	Hs.192221	ESTs	7.14
	420658	AW965215	Hs.130707	ESTs	7.12
40	408291	AB023191	Hs.44131	KIAA0974 protein	7.10
	409110	AA191493	Hs.48778	niban protein	7.10
	414485	W27026	Hs.182625	VAMP (veside-associated membrane protei	7.10
	430039	BE253012	Hs.153400	ESTs, Weakly similar to ALU1_HUMAN ALU S	7.10
45	450832	AW970602	Hs.105421	ESTs	7.10 7.08
45	417153	X57010	Hs.81343	collagen, type II, alpha 1 (primary oste	7.07
	412446	Al768015	Hs.92127	ESTS	7.06
	412953	Z45794	Hs.238809	ESTs ESTs	7.06
	418051	AW192535	Hs.19479 Hs.1395	early growth response 2 (Krox-20 (Drosop	7.04
50	421566 446999	NM_000399 AA151520	Hs.279525	hypothetical protein MGC4485	7.04
50	4 4 4 4 4 4 4	4114007010	Hs.16478	Homo sapiens cDNA: FLJ21718 fis, clone C	7.04
	440529 441111	AW207640 Al806867	Hs.126594	ESTs	7.01
	451027	AW519204	Hs.40808	ESTs -	7.00
	408432	AW195262	113.40000	gb:xn67b05.x1 NCI_CGAP_CML1 Homo sapiens	7.00
55	432223	AA333283	Hs.285336	Homo sapiens, clone IMAGE:3460280, mRNA	7.00
-	444805	AB007899	Hs.12017	homolog of yeast ubiquitin-protein ligas	6.99
	414212	AA136569	Hs.295940	KIAA0187 gene product	6.98
	431725	X65724	Hs.2839	Nome disease (pseudoglioma)	6.98
	449685	AW296669	Hs.66095	ESTs	6.97
60	447313	U92981	Hs.18081	Homo sapiens clone DT1P1B6 mRNA, CAG rep	6.96
	424590	AW966399	Hs.46821	hypothetical protein FLJ20086	6.94
	449655	AI021987	Hs.59970	ESTs	6.92
	419563	AA526235	Hs.193162	Homo sapiens cDNA FLJ11983 fis, clone HE	6.90
	434163	AW974720	Hs.25206	group XII secreted phospholipase A2	6.89
65	415809	Z32789	Hs.46601	ESTS	6.86
	425782	U66468	Hs.159525	cell growth regulatory with EF-hand doma	6.85
	417958	AA767382	Hs.193417	ESTs	6.84
	427408	AA583206	Hs.2156	RAR-related orphan receptor A	6.79
	445873	AA250970	Hs.251946	poly(A)-binding protein, cytoplasmic 1-l	6.74

	410718	Al920783	Hs.191435	ESTs	6.74
	432363	AA534489		gb:nf76g11.s1 NCL_CGAP_Co3 Homo sapiens	6.74
	436521	AW203986	Hs.213003	ESTs	6.73
~	435604	AA625279	Hs.26892	uncharacterized bone marrow protein BMO4	6.73 6.72
5	419083	AI479560	Hs.98613	Homo sapiens dDNA FLJ12292 fis, done MA	6.70
	418245	AA088767	Hs.83883	transmembrane, prostate androgen induced	6.70
	420714	BE172704	Hs.222746	KIAA1610 protein Homo sapiens cDNA: FLI21721 fis, clone C	6.67
	412707 421896	AW206373 N62293	Hs.16443 Hs.45107	ESTs	6.66
10	411078	Al222020	Hs.182364	CocoaCrisp	6.66
10	452465	AA610211	Hs.34244	ESTs	6.66
	422763	AA033699	Hs.83938	ESTs, Moderately similar to MAS2_HUMAN M	6.66
	444618	AV653785	Hs.300171	ELL-RELATED RNA POLYMERASE II, ELONGATIO	6.64
	450164	Al239923	Hs.30098	ESTs	6.63
15	431060	AF039307	Hs.249171	homeo box A11	6.62
	408031	AA081395	Hs.42173	Homo sapiens cDNA FLJ10366 fis, clone NT	6.62
	420285	AA258124	Hs.293878	ESTs, Moderately similar to ZN91_HUMAN Z	6.62
	444670	H58373	Hs.37494	hypothetical prolein MGC5370	6.62
	444489	AI151010	Hs.157774	ESTs	6.60
20	445685	AW779829	Hs.263436	gb:hn88a05.x1 NCI_CGAP_Kid11 Homo sapien	6.60
	435677	AA694142	Hs.293726	ESTs, Weakly similar to TSGA RAT TESTIS	6.59
	452221	C21322	Hs.11577	hypothetical protein FLJ22242	6.59
	431510	AA580082	Hs.112264	ESTs	6.56
05	415874	AF091622	Hs.78893	KIAA0244 protein	6.54
25	418405	AI868282	Hs.11898	ESTs, Highly similar to KIAA1370 protein	6.54
	452768	AW069459	Hs.61539	ESTs	6.54 6.52
	401451	Mocooo		ESTs	6.52
	416289	W26333 AL080276	Hs.268562	regulator of G-protein signalling 17	6.51
30	431778 409089	NM_014781	Hs.50421	KIAA0203 gene product	6.50
50	442833	AA328153	Hs.88201	ESTs, Weakly similar to A Chain A, Cryst	6.50
	431992	NM_002742	Hs.2891	protein kinase C, mu	6.49
	418833	AW974899	Hs.292776	ESTs	6.48
	429163	AA884766		gb:am20a10.s1 Soares_NFL_T_GBC_S1 Homo s	6.46
35	430403	AF039390	Hs.241382	tumor necrosis factor (ligand) superfami	6.46
	443058	AW451642	Hs.16732	ESTs	6.46
	418564	AA631143	Hs.179809	Homo sapiens prostein mRNA, complete cds	6.44
	432674	AA641092	Hs.257339	ESTs, Weakly similar to 138022 hypotheti	6.44
40	423600	A1633559	Hs.29076	ESTs	6.44
40	404253			PAT-	6.42
	433610	AA806822	Hs.112547	ESTs	6.42 6.41
	421552	AF026692	Hs.105700	secreted frizzled-related protein 4 ESTs, Weakly similar to Z223_HUMAN ZINC	6.40
	407118	AA156790 N79738	Hs.262036 Hs.136102	KIAA0853 protein	6.40
45	408608 421452	Al925946	Hs.104530	fetal hypothetical protein	6.40
43	433285	AW975944	Hs.237396	ESTs	6.40
	434926	BE543269	Hs.50252	mitochondrial ribosomal protein L32	6.40
	446189	H85224	Hs.214013	ESTs	6.40
	416806	NM_000288	Hs.79993	peroxisomal biogenesis factor 7	6.38
50	416467	H57585	Hs.37467	ESTs	6.36
-	453403	BE466639	Hs.61779	Homo sapiens cDNA FLJ13591 fis, clone PL	6.34
	429769	NM_004917	Hs.218366	kallikrein 4 (prostase, enamel matrix, p	6.34
	423642	AW452650	Hs.157148	hypothetical protein MGC13204	6.32
~ ~	425843	BE313280	Hs.159627	death associated protein 3	6.32
55	439221	AA737106	Hs.32250	ESTs, Moderately similar to 178885 serin	6.32
	428194	AA765603	Hs.180877	H3 histone, family 3B (H3.3B)	6.30
	431958	X63629	Hs.2877	cadherin 3, type 1, P-cadherin (placenta	6.30 6.30
	439366	AF100143	Hs.6540	fibroblast growth factor 13	6.30
60	452789	AW081626	Hs.242561 Hs.80247	ESTs cholecystokinin	6.30
00	416836	D54745	Hs.5364	DKFZP564I052 protein	6.29
	436962 433383	AW377314 AF034837	Hs.192731	double-stranded RNA specific adenosine d	6.29
	433383 418636	AW749855	110.102/01	gb:QV4-BT0534-281299-053-c05 BT0534 Homo	6.26
	450728	AW162923	Hs.25363	presenilin 2 (Alzheimer disease 4)	6.25
65	440293	A1004193	Hs.22123	ESTs	6.24
	453745	AA952989	Hs.63908	hypothetical protein MGC14726	6.24
	426595	AW971980	Hs.62402	p21/Cdc42/Rac1-activated kinase 1 (yeast	6.24
	444412	Al147652	Hs.216381	Homo sapiens done HH409 unknown mRNA	6.24
	413384	NM_000401	Hs.75334	exostoses (multiple) 2	6.22

		****			6.00
	426320	W47595	Hs.169300	transforming growth factor, beta 2	6.22
	423349	AF010258	Hs.127428	homeo box A9	6.20
	429165	AW009886	Hs.118258	prostate cancer associated protein 1	6.18
_	424800	AL035588	Hs.153203	MyoD family inhibitor	6.18
5	409564	AA045857	Hs.54943	fracture callus 1 (rat) homolog	6.16
	438796	W67821	Hs.109590	genethonin 1	6.16
	425451	AF242769	Hs.157461	mesenchymal stem cell protein DSC54	6.14
	451663	A1872360	Hs.209293	ESTs	6.14
	413623	AA825721	Hs.246973	ESTs.	6.12
10	452232	AW020603	Hs.271698	radial spoke protein 3	6.12
	453390	AA862496	Hs.28482	ESTs	6.12
	435542	AA687376	Hs.269533	ESTs	6.12
	420424	AB033036	Hs.97594	KIAA1210 protein	6.11
	407103	AA424881	Hs.256301	hypothetical protein MGC13170	6.10
15	409734	BE161664	Hs.56155	hypothetical protein	6.10
10	432686	BE223007	Hs.152460	Homo saplens cDNA FLJ12909 fis, clone NT	6.10
	438361	AA805666	Hs.146217	Homo sapiens cDNA: FLJ23077 fis, clone L	6.10
	411479	AW848047	1101140211	gb:IL3-CT0214-291299-052-A12 CT0214 Homo	6.10
	438849	W28948	Hs.10762	ESTs	6.08
20	452726	AF188527	Hs.61661	ESTs, Weakly similar to AF174605 1 F-box	6.08
20		D29954	Hs.13421	KIAA0056 protein	6.08
	445895	A!420611	Hs.127832	ESTs	6.07
	440774			KIAA0874 protein	6.06
	422583	- AA410506	Hs.118578	ESTs, Weakly similar to S65657 alpha-1C-	6.04
25	427500	AW970017	Hs.293948	ESTs Weakly Sittlian to 303007 appliance	6.04
23	443646	AI085198	Hs.298699	Homo sapiens cDNA FLJ13585 fis, clone PL	6.02
	410566	AA373210	Hs.43047	Homo sapiens mRNA; cDNA DKFZp586F1822 (f	6.02
	417845	AL117461	Hs.82719		6.02
	430273	Al311127	Hs.125522	ESTS	6.01
20	434792	AA649253	Hs.132458	ESTs	6.01
30	442490	AW965078	Hs.30212	thyroid receptor interacting protein 15	6.00
	420026	AI831190	Hs.166676	ESTS	6.00
	437782	Al370876	Hs.123163	exportin 1 (CRM1, yeast, homolog)	6.00
	447359	NM_012093	Hs.18268	adenylate kinase 5	6.00
25	447713	A1420733	Hs.207083	ESTs	6.00
35 ·	451073	A1758905	Hs.206063	ESTS	
•	451640	AA195601	Hs.26771	Human DNA sequence from clone 747H23 on	6.00
	410889	X91662	Hs.66744	twist (Drosophila) homolog (acrocephalos	5.97
	441222	Al277237	Hs.44208	hypothetical protein FLJ23153	5.96
40	447732	Al758398	Hs.161318	ESTs .	5.96
40	437756	AA767537	Hs.197096	ESTs	5.95
	408829	NM_006042	Hs.48384	heparan sulfate (glucosamine) 3-O-sulfot	5.94
	453911	AW503857	Hs.4007	Sarcolemmal-associated protein	5.94
	414085	AA114016	Hs.75746	aldehyde dehydrogenase 1 family, member	5.93
45	408875	NM_015434	Hs.48604	DKFZP434B168 protein	5.92
45	439451	AF086270	Hs.278554	heterochromatin-like protein 1	5.92
	423853	AB011537	Hs.133466	slit (Drosophila) homolog 1	5.91
	453060	AW294092	Hs.21594	hypothetical protein MGC15754	5.91
	420407	AA814732	Hs.145010	lipopolysaccaride-specific response 5-li	5.91
50	450480	X82125	Hs.25040	zinc finger protein 239	5.90
50	408446	AW450669	Hs.45068	hypothetical protein DKFZp434l143	5.88
	421039	NM_003478	Hs.101299	cullin 5	5.88
	451684	AF216751	Hs.26813	CDA14	5.88
	436063	AK000028	Hs.250867	ribosomal protein S24	5.86
	410507	AA355288	Hs.271408	transitional epithelia response protein	5.86
55	420179	N74530	Hs.21168	ESTs	5.84
	453878	AW964440	Hs.19025	DC32	5.84
	452270	AW975014	Hs.26	ferrochelatase (protoporphyria)	5.83
	435867	AA954229	Hs.114052	ESTs	5.82
	417683	AW566008	Hs.239154	ankyrin repeat, family A (RFXANK-like),	5.82
60	432005	AA524190	Hs.120777	ESTs, Weakly similar to ELL2_HUMAN RNA P	5.81
	406815	AA833930	Hs.288036	tRNA isopentenylpyrophosphate transferas	5.80
	437980	R50393	Hs.278436	KIAA1474 protein	5.80
	425856	AA364908	Hs.98927	hypothetical protein FLJ13993	5.79
	400301	X03635	Hs.1657	estrogen receptor 1	5.78
65	446261	AA313893	Hs.13399	hypothetical protein FLJ12615 similar to	5.78
	410141	R07775	Hs.287657	Homo sapiens cDNA: FLJ21291 fis, clone C	5.77
	427258	AA400091	Hs.39421	ESTs	5.76
	419108	AA389724	Hs.191264	ESTs, Weakly similar to ALU7_HUMAN ALU S	5.76
	442029	AW956698	Hs.14456	neural precursor cell expressed, develop	5.76
		_		•	

		•		•	
	407783	AW996872	Hs.172028	a disintegrin and metalloproteinase doma	5.75
	434408	AI031771	Hs.132586	ESTs	5.74
	415077	L41607	Hs.934	glucosaminyl (N-acetyl) transferase 2, I	5.74
_	432435	BE218886	Hs.282070	ESTs	5.74
5 -	433313	W20128	Hs.296039	ESTs	5.73
_	431740	N75450	Hs.183412	ESTs, Moderately similar to AF116721 67	5.73
			113.100412		
	412991	AW949013		gb:QV4-FT0005-110500-201-e12 FT0005 Homo	5.72
	418852	BE537037	Hs.273294	hypothetical protein FLJ20069	5.72
	418882	NM_004996	Hs.89433	ATP-binding cassette, sub-family C (CFTR	5.72
10					5.72
10	446867	AB007891	Hs.16349	KIAA0431 protein	
	437866	AA156781	Hs.83992	metallothionein 1E (functional)	5.72
	410232	AW372451	Hs.61184	CGI-79 protein	5.70
	414452	AA454038	Hs.29032	ESTs	5.70
	422762	AL031320	Hs.119976	Human DNA sequence from clone RP1-20N2 o	5.70
15	428730	AA625947	Hs.25750	ESTs	5.70
	431571	AW500486	Hs.180610	splicing factor proline/glutamine rich (5.70
			Hs.98074	itchy (mouse homolog) E3 ubiquitin prote	5.70
	433393	AF038564			
	450616	AL133067	Hs.25214	hypothetical protein	5.70
	443774	AL117428	Hs.9740	DKFZP434A236 protein	5.69
20	446100	AW967109	Hs.13804	hypothetical protein dJ462023.2	5.69
20					5.68
	419168	Al336132	Hs.33718	Homo sapiens cDNA FLJ12641 fis, clone NT	
	416653	AA768553	Hs.77496	metallothionein 1E (functional)	5.67
	452679	Z42387	Hs.4299	transmembrane, prostate androgen induced	5.66
	450244	AA007534	Hs.125062	ESTs	5.66
25			Hs.46638		5.65
23	408621	A1970672		chromosome 11 open reading frame 8	
	450325	Al935962	Hs.26289	ESTs	5.65
	439671	AW162840	Hs.6641	kinesin family member 5C	5.64
	452387	Al680772	Hs.4316	trinucleotide repeat containing 12	5.64
	413992	W26276	Hs.136075	RNA, U2 small nuclear	5.63
30	444151	AW972917	Hs.128749	alpha-methylacyl-CoA racemase	5.63
	417791	AW965339	Hs.111471	ESTs	5.62
		Al936442	Hs.59838	hypothetical protein FLJ10808	5.60
	410196		112533000		
	415123	D60925		ESTs	5.60
	429170	NM_001394	Hs.2359	dual specificity phosphatase 4	5.60
35	434415	BE177494		gb:RC6-HT0596-270300-011-C05 HT0596 Homo	5.60
	440738	Al004650	Hs.225674	WD repeat domain 9	5.60
	443830	Al142095	Hs.143273	ESTs	5.60
	449603	Al655662	Hs.197698	ESTs	5.60
	414342	AA742181	Hs.75912	KIAA0257 protein	5.59
40	422634	NM_016010	Hs.118821	CGI-62 protein	5.56
70		_			
	435047	AA454985	Hs.54973	cadherin-like protein VR20	5.55
	400268				5.55
	452055	Al377431	Hs.293772	hypothetical protein MGC10858	5.54
	437073	Al885608	Hs.94122	ESTs	5.54
15					
45	434072	H70854	Hs.283059	Homo sapiens PRO1082 mRNA, complete cds	5.53
	418339	AA639902	Hs.104215	ESTs, Moderately similar to SPCN_HUMAN S	5.52
	434551	BE387162	Hs.280858	ESTs, Highly similar to A35661 DNA excis	5.52
	439569	AW602166	Hs.222399	CEGP1 protein	5.51
ec.	441102	AA973905	Hs.16003	intermediate filament protein syncoilin	5.50
50	448310	Al480316		gb:tm26h09,x1 Soares_NFL_T_GBC_S1 Homo s	5.50
	413173	BE076928	Hs.70980	ESTs	5.48
	436246	AW450963	Hs.119991	ESTs	5.48
	449300	Al656959	Hs.222165	ESTs	5.48
	452823	AB012124	Hs.30696	transcription factor-like 5 (basic helix	5.48
55	451403	AA885569	Hs.15727	Homo sapiens cDNA FLJ14511 fis, clone NT	5.46
	417061	Al675944	Hs.188691	Homo sapiens cDNA FLJ12033 fis, clone HE	5.44
	429126	AW172356	Hs.99083	ESTs	5.44
	431316	AA502663	Hs.145037	ESTs	5.44
	439192	AW970536	Hs.105413	ESTs	5.44
60				specific granule protein (28 kDa); cyste	5.44
50	431938	AA938471	Hs.115242		
	451552	AA047233	Hs.33810	ESTs	5.43
	416991	N36389	Hs.295091	KIAA0226 gene product	5.42
	427638	AA406411	Hs.208341	ESTs, Weakly similar to KIAA0989 protein	5.42
				ESTs	5.42
65	427718	AI798680	Hs.25933		
65	438710	AA833907	Hs.178724	ESTs, Weakly similar to ALU1_HUMAN ALU S	5.42
	406076	AL390179	Hs.137011	Homo sapiens mRNA; cDNA DKFZp547P134 (fr	5.40
	431263	AW129203	Hs.13743	ESTs	5.40
			Hs.103042	microtubule-associated protein 1B	5.38
	421264	AL039123			
	421685	AF189723	Hs.106778	ATPase, Ca++ transporting, type 2C, memb	5.37

	408460	AA054726	Hs.285574	ESTs	5.36
	409091	AW970386	Hs.269423	ESTs	5.36
	421987	Al133161 .	Hs.286131	CGI-101 protein	5.36
_	428002	AA418703		gb:zv98c03.s1 Soares_NhHMPu_S1 Homo sapi	5.36
5	441217	Al922183	Hs.213246	ESTs	5.36
	426006	R49031	Hs.22627	ESTs	5.35
	422806	BE314767	Hs.1581	glutathione S-transferase theta 2	5.34 5.32
	432281 451982	AK001239 F13038	Hs.274263 Hs.27373	hypothetical protein FLJ10377 Homo saplens mRNA; cDNA DKFZp56401763 (f	5.32
10	421129	BE439899	Hs.89271	ESTs	5.31
10	444042	NM_004915	Hs.10237	ATP-binding cassette, sub-family G (WHIT	5.31
	410150	AW382942	Hs.6774	ESTs	5.30
	423952	AW877787	Hs.136102	KIAA0853 protein	5.30
	452822	X85689	Hs.288617	hypothetical protein FLJ22621	5.30
15	447752	M73700	Hs.347	lactotransferrin	5.29
	441766	R53790	Hs.23294	hypothetical protein FLJ14393	5.29 5.27
	431359 427212	AW993522 AW293849	Hs.292934 Hs.58279	ESTs ESTs, Weakly similar to ALU7_HUMAN ALU S	5.27 5.27
	449916	T60525	Hs.299221	pyruvate dehydrogenase kinase, isoenzyme	5.27
20	454014	AW016670	Hs.233275	ESTs	5.27
	419714	AA758751	Hs.98216	ESTs	5.26
	428845	AL157579	Hs.153610	KIAA0751 gene product	5.26
	417333	AL157545	Hs.42179	bromodomain and PHD finger containing, 3	5.24
25	419986	Al345455	Hs.78915	GA-binding protein transcription factor,	5.24
25	407182	AA312551	Hs.230157	ESTS	5.22 5.22
	420111 428058	AA255652 AI821625	Hs.191602	gb:zs21h11.r1 NCI_CGAP_GCB1 Homo saplens ESTs	5.22
	459551	AI472808	113.131002	gb:tj70e07.x1 Soares_NSF_F8_9W_OT_PA_P_S	5.22
	432524	Al458020	Hs.293287	ESTs	5.22
30	436207	AA334774	Hs.12845	hypothetical protein MGC13159	5.22
	410870	U81599	Hs.66731	homeo box B13	5.22
	451418	BE387790	Hs.26369	hypothetical protein FLJ20287	5.22
	409757	NM_001898	Hs.123114	cystatin SN	5.21 5.21
35	441124 428593	T97717 AW207440	Hs.119563 Hs.185973	ESTs degenerative spermatocyte (homolog Droso	5.21
55	436401	A1087958	Hs.29088	ESTs	5.20
	437113	AA744693	1.0.2000	gb:ny26c10.s1 NCI_CGAP_GCB1 Homo sapiens	5.20
	450947	Al745400	Hs.204662	ESTS	5.20
40	453279	AW893940	Hs.59698	ESTs	5.20
40	445467	AI239832	Hs.15617	ESTs, Weakly similar to ALU4_HUMAN ALU S	5.19
	448944	AB014605	Hs.22599	atrophin-1 interacting protein 1; activi	5.19 5.18
	412198 422646	AA937111 H87863	Hs.69165 Hs.151380	ESTs ESTs, Weakly similar to T16584 hypotheti	5.18
	438986	AF085888	Hs.269307	ESTs	5.18
45	453954	AW118336	Hs.75251	DEAD/H (Asp-Glu-Ala-Asp/His) box binding	5.18
	447541	AK000288	Hs.18800	hypothetical protein FLJ20281	5.18
	434029	AA621763	Hs.170434	Homo sapiens cDNA FLJ14242 fis, clone OV	5.16
	459294	AW977286	Hs.169531	RBP1-like protein	5.16
50	429441 424692	AJ224172 AA429834	Hs.204096 Hs.151791	lipophilin B (uteroglobin family member) KIAA0092 gene product	5.16 5.15
50	427359	AW020782	Hs.79881	Homo sapiens cDNA: FLI23006 fis, clone L	5.15
	419872	A1422951	Hs.146162	ESTs	5.15
	429422	AK001494	Hs.202596	Homo sapiens cDNA FLJ10632 fis, cloné NT	5.14
	448902	Z45998	Hs.22543	Homo sapiens mRNA; cDNA DKFZp761l1912 (f	5.14
55	459055	N23235	Hs.30567	ESTs, Weakly similar to B34087 hypotheti	5.14
	431318	AA502700	Hs.293147	ESTs, Moderately similar to A46010 X-lin	5.14
	452953	Al932884	Hs.271741	ESTs, Weakly similar to A46010 X-linked hypothetical protein FLJ22104	5.13 5.12
	428372 434401	AK000684 Al864131	Hs.183887 Hs.71119	Putative prostate cancer tumor suppresso	5.12
60	416434	AW163045	Hs.79334	nuclear factor, interleukin 3 regulated	5.11
	410268	AA316181	Hs.61635	six transmembrane epithelial antigen of	5.10
	417517	AF001176	Hs.82238	POP4 (processing of precursor, S. cerev	5.10
	453616	NM_003462	Hs.33846	dynein, axonemal, light intermediate pol	5.10
45	427958	AA418000	Hs.98280	potassium intermediate/small conductance	5.09
65	407945	X69208	H\$.606	ATPase, Cu++ transporting, alpha polypep collagen, type IX, alpha 1	5.08
	425154 412863	NM_001851 AA121673	Hs.154850 Hs.59757	zinc finger protein 281	5.08 5.06
	420807	AA280627	Hs.57846	ESTs	5.06
	430568	AA769221	Hs.270847	delta-tubulin	5.06

		4.7740004			E 00
	433687	AA743991		gb:ny57g01.s1 NCL_CGAP_Pr18 Homo sapiens	5.06
	438375	AW015940	Hs.232234	ESTs	5.06
	418092	R45154	Hs.106604	ESTs	5.06
	418576	AW968159	Hs.289104	Alu-binding protein with zinc finger dom	5.05
5	413328	Y15723	Hs.75295	guanylate cyclase 1, soluble, alpha 3	5.04
_	414271	AK000275	Hs.75871	protein kinase C binding protein 1	5.04
	432729	AK000292	Hs.278732	hypothetical protein FLJ20285	5.04
	433433	Al692623	Hs.121513	Homo sapiens clone Z'3-1 placenta expres	5.04
	439662		Hs.269060	ESTs	5.04
10		H97552			5.04
10	439743	AL389956	Hs.283858	Homo sapiens mRNA full length insert cDN	
	417511	AL049176	Hs.82223	chordin-like	5.02
	437814	Al088192	Hs.135474	ESTs, Weakly similar to DDX9_HUMAN ATP-D	5.02
	426342	AF093419	Hs.169378	multiple PDZ domain protein	5.02
	429782	NM_005754	Hs.220689	Ras-GTPase-activating protein SH3-domain	5.02
15	429975	Al167145	Hs.165538	ESTs	5.02
	436209	AW850417	Hs.254020	ESTs, Moderately similar to unnamed prot	5.02
	438571	AW020775	Hs.56022	ESTs	5.02
	450223	AA418204	Hs.241493	natural killer-tumor recognition sequenc	5.02
			Hs.267705	tubulin-specific chaperone e	5.01
20	408267	AW380525	113.207700	gb:HSC28F061 normalized infant brain cDN	5.00
20	417730	Z44761	11-4004		
	425465	L18964	Hs.1904	protein kinase C, iota	5.00
	430599	NM_004855	Hs.247118	phosphatidylinositol glycan, class B	5.00
	450961	AW978813	Hs.250867	metallothionein 1E (functional)	5.00
	451386	AB029006	Hs.26334	spastic paraplegia 4 (autosomal dominant	5.00
25	420380	AA640891	Hs.102406	ESTs	4.99
	424947	R77952	Hs.239625	ESTs, Weakly similar to alternatively sp	4.99
	442653	BE269247	Hs.170226	gb:601185486F1 NIH_MGC_8 Homo sapiens cD	4.98
	457211	AW972565	Hs.32399	ESTs, Wealdy similar to S51797 vasodilat	4.97
	425851	NM_001490	Hs.159642	glucosaminyl (N-acetyl) transferase 1, c	4.97
30		AA490770	Hs.182382	ESTs	4.96
30	446279			ESTs	4.96
	433377	Al752713	Hs.43845		4.96
	450218	R02018	Hs.168640	ankylosis, progressive (mouse) homolog	
	412715	NM_000947	Hs.74519	primase, polypeptide 2A (58kD)	4.94
~=	- 448164	R61680	Hs.26904	ESTs, Moderately similar to Z195_HUMAN Z	4.94
35	420121	AW968271	Hs.191534	ESTs, Weakly similar to ALU1_HUMAN ALU S	4.94
	421689	N87820	Hs.106826	KIAA1696 protein	4.93
	445808	AV655234	Hs.298083	ESTs, Moderately similar to PC4259 ferri	4.92
	416533	BE244053	Hs.79362	retinoblastoma-like 2 (p130)	4.92
	418049	AA211467	Hs.190488	Homo sapiens, Similar to nuclear localiz	4.92
40	436039	AW023323	Hs.121070	ESTs	4.92
	432653	N62096	Hs.293185	ESTs, Weakly similar to JC7328 amino aci	4.91
	420324	AF163474	Hs.96744	prostate androgen-regulated transcript 1	4.91
	403047	74 100414	113.001 44	products unaregen regulates summarks	4.91
		A A7040E0	No 201567	ESTs	4.90
15	436899	AA764852	Hs.291567	T-11.	4.90
45	431117	AF003522	Hs.250500	delta (Drosophila)-like 1	
	427617	D42063	Hs.179825	RAN binding protein 2	4.88
	428804	AK000713	Hs.193736	hypothetical protein FLJ20706	4.88
	433050	Al093930	Hs.163440	Homo sapiens cDNA: FLJ21000 fis, clone C	4.88
	418575	AA225313	Hs.222886	ESTs, Weakly similar to TRHY_HUMAN TRICH	4.86
50	432615	AA557191	Hs.55028	ESTs, Weakly similar to I54374 gene NF2	4.86
	412652	AI801777	Hs.6774	ESTs	4.86
	432473	Al202703	Hs.152414	ESTs	4.86
	449071	NM_005872	Hs.22960	breast carcinoma amplified sequence 2	4.86
	450654	AJ245587	Hs.25275	Kruppel-type zinc finger protein	4.85
55	418866	T65754	Hs.100489	gb:yc11c07.s1 Stratagene lung (937210) H	4.85
33			113.100-103	gb:yg30f05.r1 Soares fetal liver spleen	4.84
	407596	R86913	LI- 00074C		4.84
	456516	BE172704	Hs.222746	KIAA1610 protein	
	426501	AW043782	Hs.293616	ESTs	4.84
<i></i>	448730	AB032983	Hs.21894	KIAA1157 protein	4.84
60	458339	AW976853	Hs.172843	ESTs	4.83
	422083	NM_001141	Hs.111256	arachidonate 15-lipoxygenase, second typ	4.82
	420159	AI572490	Hs.99785	Homo saplens cDNA: FLJ21245 fis, clone C	4.82
	424103	NM_001918	Hs.139410	dihydrolipoamide branched chain transacy	4.82
	449535	W15267	Hs.23672	low density lipoprotein receptor-related	4.82
65	422048	NM_012445	Hs.288126	spondin 2, extracellular matrix protein	4.82
-	416737	AF154335	Hs.79691	LIM domain protein	4.82
				golgin-67	4.81
	419972	AL041465	Hs.294038		4.81
	420235	AA256756	Hs.31178	ESTs	
	423412	AF109300	Hs.147924	prostate cancer associated protein 5	4.80

				•	
	429598	AA811257	Hs.269710	ESTs	4.80
	457114	A1821625	Hs.191602	ESTs	4.80
	421828	AW891965	Hs.289109	histone deacetylase 3	4.79
5	424602	AK002055	Hs.301129	hypothetical protein FLJ11193	4.78
2	428364	AA426565	Hs.160541	ESTs, Moderately similar to ALU1_HUMAN A	4.78
	452335 410765	AW188944 Al694972	Hs.61272 Hs.66180	ESTs nucleosome assembly protein 1-like 2	4.78 4.77
	421040	AA715026	Hs.135280	ESTs	4.76
	421518	AI056392	Hs.208819	ESTs	4.76
10	452560	BE077084	113200010	ESTs	4.76
	409752	AW963990		gb:EST376063 MAGE resequences, MAGH Homo	4.75
	439703	AF086538	Hs.196245	ESTs	4.75
	418836	Al655499	Hs.161712	ESTs	4.74
	450642	R39773	Hs.7130	copine IV	4.74
15	419879	Z17805	Hs.93564	Homer, neuronal immediate early gene, 2	4.74
	411440	AW749402		gb:QV4-BT0383-281299-061-c06 BT0383 Homo	4.74
	450649	NM_001429	Hs.297722	E1A binding protein p300	4.74
	408738	NM_014785	Hs.47313	KIAA0258 gene product	4.73
20	435020	AW505076	Hs.301855	DiGeorge syndrome critical region gene 8	4.72 4.72
20	411624 439360	BE145964 AA448488	Hs.55346	KIAA0594 protein ribosomal protein L44	4.72
	440491	R35252	Hs.24944	ESTs, Weakly similar to 2109260A B cell	4.72
	442611	BE077155	Hs.177537	hypothetical protein DKFZp761B1514	4.72
	443555	N71710	Hs.21398	ESTs, Moderately similar to A Chain A, H	4.72
25	453800	BE300741	Hs.288416	hypothetical protein FLJ13340	4.72
	457528	AW973791	Hs.292784	ESTs	4.72
	416795	Al497778	Hs.168053	HBV pX associated protein-8	4.71
	407302	R74206	Hs.268755	ESTs, Weakly similar to I78885 serine/th	4.71
30	404721	414040040	11- 400070		4.70
30	426261	AW242243 AK000850	Hs.168670	peroxisomal famesylated protein Homo sapiens cDNA FLJ20843 fis, clone AD	4.70 4.70
	431924 435256	AF193766	Hs.272203 Hs.13872	cytokine-like protein C17	4.70
	438295	Al394151	Hs.37932	ESTs	4.70
	442655	AW027457	Hs.30323	ESTs, Weakly similar to B34087 hypotheti	4.70
35	415788	AW628686	Hs.78851	KIAA0217 protein	4.69
	442760	BE075297	Hs.10067	ESTs, Weakly similar to A43932 mucin 2 p	4.69
	432432	AA541323	Hs.115831	ESTs	4.68
	454398	AA463437	Hs.11556	Homo sapiens cDNA FLJ12566 fis, clone NT	4.68
40	452741	BE392914	Hs.30503	Homo sapiens cDNA FLJ11344 fis, clone PL	4.67
40	424853	BE549737	Hs.132967	Human EST clone 122887 mariner transposo	4.67 4.66
	419706 412088	C04649 A1689496	Hs.77899 Hs.108932	tropomyosin 1 (alpha) ESTs	4.65
	416276	U41060	Hs.79136	LIV-1 protein, estrogen regulated	4.64
	429281	AA830856	Hs.29808	Homo sapiens cDNA: FLJ21122 fis, clone C	4.64
45	448207	A1475490	Hs.170577	ESTs	4.64
	408374	AW025430	Hs.155591	forkhead box F1	4.64
	447162	BE328091	Hs.157396	ESTs, Weakly similar to A46010 X-linked	4.64
	451900	AB023199	Hs.27207	KIAA0982 protein	4.63
50	421437	AW821252	Hs.104336	hypothetical protein	4.63
30	418624	A1734080	Hs.104211	ESTs	4.63
	426172	AA371307 AW136488	Hs.125056	ESTs ESTs	4.62 4.61
	439831 452994	AW962597	Hs.25545 Hs.31305	KIAA1547 protein	4.61
	457726	Al217477	Hs.194591	ESTs	4.60
55	434629	AA789081	Hs.4029	glioma-amplified sequence-41	4.60
	403764			• • • • • • • • • • • • • • • • • • • •	4.58
	410659	Al080175	Hs.68826	ESTs	4.58
	432383	AK000144	Hs.274449	Homo sapiens cDNA FLJ20137 fis, clone CO	4.58
4 0	451246	AW189232	Hs.39140	cutaneous T-cell lymphoma tumor antigen	4.58
60	433234	AB040928	Hs.65366	KIAA1495 protein	4.57
	424983	A1742434	Hs.169911	ESTs ESTs Wooldy similar to OAU ID1 debdoors:	4.56
	437812	AI582291	Hs.16846	ESTs, Weakly similar to O4HUD1 debrisoqu hypothetical protein FLJ13409; KIAA1711	4.56 4.55
	438447 434715	A1082883 BE005346	Hs.167593 Hs.116410	ESTs	4.55
65	447673	A!823987	Hs.182285	ESTS	4.54
J J	408897	N50204	Hs.283709	lipopolysaccharide specific response-7 p	4.54
	436645	AW023424	Hs.156520	ESTs	4.54
	421247	BE391727	Hs.102910	general transcription factor IIH, polype	4.53
	450377	AB033091	Hs.24936	KIAA1265 protein	4.53

					4.00
	433644	AW342028	Hs.256112	gb:hb75d03.x1 NCI_CGAP_Ut2 Homo saplens	4.53
	408321	AW405882	Hs.44205	cortistatin	4.53
	439225	AA192669	Hs.45032	ESTs	4.52
	440348	AW015802	Hs.47023	ESTs	4.52
5					4.52
J	446351	AW444551	Hs.258532	x 001 protein	
	451212	AW902672	Hs.287334	ESTs	4.52
	430294	AI538226	Hs.135184	guanine nucleotide binding protein 4	4.52
	435005	U80743	Hs.4316	trinucleotide repeat containing 12	4.52
	448072	Al459306	Hs.24908	ESTs	4.50
10		M403000	113.24300	LOIS	4.50
10	403721				
	451018	AW965599	Hs.247324	mitochondrial ribosomal protein S14	4.50
	453070	AK001465	Hs.31575	SEC63, endoplasmic reticulum translocon	4.49
	417412	X16896	Hs.82112	interleukin 1 receptor, type i	4.48
	439735	A1635386	Hs.142846	hypothetical protein	4.48
15					4.48
15	435663	AI023707	Hs.134273	ESTs	
	424036	AA770688	Hs.81946	H2A histone family, member L	4.48
	426386	AA748850	Hs.174877	bladder cancer overexpressed protein	4.48
	408622	AA056060	Hs.202577	Homo sapiens cDNA FLJ12166 fis, clone MA	4.47
	444269	AJ590346	Hs.146220	ESTs	4.47
20	430187	Al799909	Hs.158989	ESTs	4.46
20					
	427761	AA412205	Hs.140996	ESTs	4.46
	430261	AA305127	Hs.237225	hypothetical protein HT023	4.46
	444169	AV648170	Hs.58756	ESTs	4.44
	430598	AK001764	Hs.247112	hypothetical protein FLJ10902	4.44
25	412903	BE007967	Hs.155795	ESTs	4.44
23				=	4.44
	417048	A1088775	Hs.55498	geranylgeranyl diphosphate synthase 1	
	442710	AI015631	Hs.23210	ESTs	4.44
	457413	AA743462	Hs.165337	ESTs	4.44
	400303	AA242758	Hs.79136	LIV-1 protein, estrogen regulated	4.42
30	443268	AI800271	Hs.129445	hypothetical protein FLJ12496	4.42
50		AL120659	Hs.6111	aryl-hydrocarbon receptor nuclear transl	4.42
	438209			·_ ·	4.41
	431724	AA514535	Hs.283704	ESTs	
	412280	AW205116	Hs.272814	hypothetical protein DKFZp434E1723	4.40
	440801	AA906366	Hs.190535	ESTs	4.40
35	452959	AI933416	Hs.189674	ESTs	4.40
	453861	AI026838	Hs.30120	ESTs, Weakly similar to NUCL_HUMAN NUCLE	4.40
				nuclear receptor subfamily 4, group A, m	4.40
	417421	AL138201	Hs.82120		
	447270	AC002551	Hs.331	general transcription factor IIIC, polyp	4.38
	433641	AF080229		gb:Human endogenous retrovirus K clone 1	4.38
40	447078	AW885727	Hs.301570	ESTs .	4.38
	424242	AA337476		hypothetical protein MGC13102	4.37
	408170	AW204516	Hs.31835	ESTs	4.36
				TATA box binding protein (TBP)-associate	4.36
	448757	Al366784	Hs.48820		
	420021	AA252848	Hs.293557	ESTs	4.36
45	449694	A1659790	Hs.253302	ESTs	4.36
	453867	Ai929383	Hs.108196	hypothetical protein DKFZp434N185	4.36
	458712	Al347502	Hs.173066	hypothetical protein FLJ20761	4.36
				ESTs, Weakly similar to YK54_YEAST HYPOT	4.35
	417251	AW015242	Hs.99488		4.35
50	434423	NM_006769	Hs.3844	LIM domain only 4	
50	423427	AL137612	Hs.285848	KIAA1454 protein	4.34
	415715	F30364		ESTs	4.33
	404561				4.32
	422969	AA782536	Hs.122647	N-myristoyltransferase 2 -	4.32
	423685	BE350494	Hs.49753	uveal autoantigen with coiled coil domai	4.32
55				ESTs, Weakly similar to I38022 hypotheti	4.32
23	443977	AL120986	Hs.150627		
	425071	NM_013989	Hs.154424	deiodinase, lodothyronine, type II	4.32
	431583	AL042613	Hs.262476	S-adenosylmethionine decarboxylase 1	4.31
	411379	AI816344	Hs.12554	ESTs, Weakly similar to NPL4_HUMAN NUCLE	4.30
	421476	AW953805	Hs.21887	ESTs	4.30
60		H16097		ESTs	4.30
UU	425178		Hs.161027		
	439262	AA832333	Hs.124399	ESTs	4.30
	442818	AK001741	Hs.8739	hypothetical protein FLJ10879	4.30
	421977	W94197	Hs.110165	ribosomal protein L26 homolog	4.29
	437114	AA836641	Hs.163085	ESTs	4.28
65	420195	N44348	Hs.300794	Homo sapiens cDNA FLJ11177 fis, clone PL	4.28
0.5				ESTs .	4.27
	418330	BE409405	Hs.94722		
	419750	AL079741	Hs.183114	Homo sapiens cDNA FLJ14236 fis, done NT	4.26
	437065	AL036450	Hs.103238	ESTs	4.26
	455276	BE176479		gb:RC3-HT0585-160300-022-b09 HT0585 Homo	4.24
				-	

	416292	AA179233	Hs.42390	nasopharyngeal carcinoma susceptibility	4.24
	423740	Y07701	Hs.132243	aminopeptidase puromycin sensitive	4.24
	442023	Al187878	Hs.144549	ESTs	4.24
_	426764	AA732524	Hs.151464	ESTs, Weakly similar to ALUC_HUMAN IIII	4.23
5	454058	Al273419	Hs.135146	hypothetical protein FLJ13984	4.23
	456511	AA282330	Hs.145668	ESTs	4.22
**	448330	AL036449	Hs.207163	ESTs	4.22
	424701	NM_005923	Hs.151988	mitogen-activated protein kinase kinase	4.21
10	432621	Al298501	Hs.12807	ESTs, Weakly similar to T46428 hypotheti	4.20
10	445707 419910	Al248720 AA662913	Hs.114390 Hs.190173	ESTs ESTs, Weakly similar to A46010 X-linked	4.20 4.20
	424085	NM 002914	Hs.139226	replication factor C (activator 1) 2 (40	4.20
	440749	W22335	Hs.7392	hypothetical protein MGC3199	4.20
	442787	W93048	Hs.227203	hypothetical protein MGC2747	4.20
15	443414	R54594	Hs.25209	ESTs	4.20
	443556	AA256769	Hs.94949	methylmalonyl-CoA epimerase	4.20
	444170	AW613879	Hs.102408	ESTs	4.20
	446751	AA766998	Hs.85874	Human DNA sequence from clone RP11-16L21	4.20
20	421041	N36914	Hs.14691	ESTs, Moderately similar to 138022 hypot	4.19
20 -	447476	BE293466	Hs.20880	ESTs, Weakly similar to 138022 hypotheti	4.19
•	448543	AW897741 AB014515	Hs.21380 Hs.288891	Homo sapiens mRNA; cDNA DKFZp586P1124 (f KIAA0615 gene product	-4.18 4.18
	410294 433607	AA602004	Hs.23260	ESTs	4.18
	435552	Al668636	Hs.193480	ESTs, Moderately similar to ALU6_HUMAN A	4.18
25	447124	AW976438	Hs.17428	RBP1-like protein	4.18
	453308	AW959731	Hs.32538	ESTs	4.17
	439328	W07411	Hs.118212	ESTs, Moderately similar to ALU3_HUMAN A	4.16
	430473	AW130690	Hs.299842	ESTs	4.16
20	437257	Al283085	Hs.290931	ESTs, Weakly similar to YFJ7_YEAST HYPOT	4.16
30	438018	AK001160	Hs.5999	hypothetical protein FLJ10298	4.16
	443857	AI089292	Hs.287621	hypothetical protein FLJ14069	4.15
	446711 419103	AF169692 Z40229	Hs.12450 Hs.96423	protocadherin 9	4.15 4.14
	405403	240229	NS.30423	hypothetical protein FLJ23033	4.14
35	407378	AA299264		ESTs, Moderately similar to I38022 hypot	4.14
	408986	AW298602	Hs.197687	ESTs	4.14
	418727	AA227609	Hs.94834	ESTs	4.14
	434400	Al478211	Hs.186896	Homo sapiens cDNA FLJ11417 fis, clone HE	4.14
40	438578	AA811244	Hs.164168	ESTs	4.14
40	450459	Al697193	Hs.299254	Homo sapiens cDNA: FLJ23597 fis, clone L	4.14
	429887	AW366286	Hs.145696	splicing factor (CC1.3)	4.13
	448148	NM_016578	Hs.20509 Hs.17850	HBV pX associated protein-8	4.13 4.12
	450316 417531	W84446 NM_003157	Hs.1087	hypothetical protein MGC4643 serine/threonine kinase 2	4.12
45	431592	R69016	Hs.293871	hypothetical protein MGC10895s	4.12
-	432463	AA548518	Hs.186733	ESTs	4.12
	433613	AA836126	Hs.5669	ESTs	4.12
	434739	AA804487	Hs.144130	ESTs	4.12
٠.	438259	AW205969	Hs.131808	ESTs	4.12
50	425810	Al923627	Hs.31903	ESTs	4.10
	432672	AW973775	Hs.130760	myosin phosphatase, target subunit 2	4.10
	433345	AI681545	Hs.152982 Hs.288031	hypothetical protein FLJ13117 sterol-C5-desaturase (fungal ERG3, delta	4.10 4.09
	432712 453020	AB016247 AL162039	Hs.31422	Homo saplens mRNA; cDNA DKFZp434M229 (fr	4.09
55	412045	AA099802	Hs.4299	transmembrane, prostate androgen induced	4.09
	435114	AA775483	Hs.288936	mitochondrial ribosomal protein L9	4.08
	443204	AW205878	Hs.29643	Homo saplens cDNA FLJ13103 fis, clone NT	4.08
	445459	Al478629	Hs.158465	likely ortholog of mouse putative IKK re	4.08
CO	438938	H46212	Hs.137221	ESTs	4.07
60	454119	BE549773	Hs.40510	uncoupling protein 4	4.06
	411000	N40449	Hs.201619	ESTs, Weakly similar to S38383 SEB4B pro	4.06
	418926	AA232658	Hs.87070	UDP-glucose:glycoprotein glucosyltransfe	4.06
	424432	AB037821 AA002064	Hs.146858	protocadherin 10	4.06
65	449673 429299	AAUU2U64 A1620463	Hs.18920 Hs.99197	ESTs hypothetical protein MGC13102	4.06 4.06
J	429299 422174	AL049325	Hs.112493	Homo sapiens mRNA; cDNA DKFZp564D036 (fr	4.05
	455497	AA112573	Hs.285691	Homo sapiens prostein mRNA, complete cds	4.05
	415138	C18356	Hs.78045	tissue factor pathway Inhibitor 2	4.04
	402791			• • •	4.04

	426792	AL044854	Hs.172329	KIAA0576 protein	4.04
	438660	U95740	Hs.6349	Homo sapiens, clone IMAGE:3010666, mRNA,	4.04
	442768	AL048534	Hs.48458	ESTs, Weakly similar to ALU8_HUMAN ALU S	4.04
	447568	AF155655	Hs.18885		4.04
5				CGI-116 protein	
J	428342	Al739168	Hs.131798	Homo sapiens cDNA FLJ13458 fis, clone PL	4.04
	453439	Al572438	Hs.32976	guanine nucleotide binding protein 4	4.02
	453857	AL080235	Hs.35861	DKFZP586E1621 protein	4.02
	428249	AA130914	Hs.183291	zinc finger protein 268	4.02
	432015	AL157504	Hs.159115	Homo sapiens mRNA; cDNA DKFZp586O0724 (f	4.02
10					4.02
10	445495	BE622641	Hs.38489	ESTs, Weakly similar to 138022 hypotheti	
	451746	M86178		ESTs	4.02
	452211	Al985513	Hs.233420	ESTs	4.02
	453046	AA284040	Hs.219441	ESTs, Highly similar to CA5B_HUMAN CARBO	4.02
	456038	AA203285	Hs.294141	ESTs, Weakly similar to alternatively sp	4.02
15	452449	AW068658	Hs.20943	ESTs	4.02
13					
	407204	R41933	Hs.140237	ESTs, Weakly similar to ALU1_HUMAN ALU S	4.01
	428046	AW812795	Hs.155381	ESTs, Moderately similar to 138022 hypot	4.01
	438520	AA706319	Hs.98416	ESTs	4.01
	443292	AK000213	Hs.9196	hypothetical protein	4.01
20	432715	AA247152	Hs.200483	ESTs, Weakly similar to KIAA1074 protein	4.00
	403797		1.0400.00	and the first of t	4.00
		A A O 4 C 4 4 O	U- 000000	about Contract NO. OOAD Det Home contract	4.00
	418347	AA216419	Hs.269295	gb:nc16e03.s1 NCI_CGAP_Pr1 Homo saplens	*
	419459	AW291128	Hs.278422	DKFZP586G1122 protein	4.00
	420911	U77413	Hs.100293	O-linked N-acetylglucosamine (GlcNAc) tr	4.00
25	425176	AW015644	Hs.301430	TEA domain family member 1 (SV40 transcr	4.00
	447505	AL049266	Hs.18724	Homo sapiens mRNA; cDNA DKFZp564F093 (fr	4.00
		AL133761	110.10.2	gb:DKFZp761C1413_r1 761 (synonym: hamy2)	4.00
	453773		LI= 400040		
	434384	AA631910	Hs.162849	ESTs	3.99
	422471	AA311027	Hs.271894	ESTs, Weakly similar to I38022 hypotheti	3.99
30	427386	AW836261	Hs.177486	ESTs	3.98
	433394	Al907753	Hs.93810	cerebral cavemous malformations 1	3.98
	441269	AW015206	Hs.178784	ESTs	3.97
		AB020695	Hs.91662	KIAA0888 protein	3.96
	419629				
25	435008	AF150262	Hs.162898	ESTs	3.96
35	456649	R74441	Hs.117176	poly(A)-binding protein, nuclear 1	3.96
	418723	AA504428	Hs.10487	Homo sapiens, clone IMAGE:3954132, mRNA,	3.96
	428738	NM_000380	Hs.192803	xeroderma pigmentosum, complementation g	3.95
	430456	AA314998	Hs.241503	hypothetical protein	3.95
	422017	NM_003877	Hs.110776	STAT induced STAT inhibitor-2	3.95
40					
40	409960	BE261944	Hs.153028	hexokinase 1	3.95
	455309	AW894017		gb:RC4-NN0027-150400-012-g04 NN0027 Homo	3.95
	450295	A1766732	Hs.201194	ESTs .	3.94
	456660	AA909249	Hs.112282	solute carrier family 30 (zinc transport	3.94
	410908	AA121686	Hs.10592	ESTs	3.94
45	447145	AA761073	Hs.192943	TRAF family member-associated NFKB activ	3.94
73					
	449318	AW236021	Hs.108788	Homo sapiens, Similar to RIKEN cDNA 5730	3.94
	449869	W57990	Hs.60059	Homo sapiens cDNA FLJ11478 fis, clone HE	3.94
	411887	AW182924	Hs.128790	ESTs	3.93
	437531	Al400752 ·	Hs.112259	T cell receptor gamma locus	3.93
50	452238	F01811	Hs.187931	ESTs	3.93
	410486	AW235094	Hs.193424	zinc finger protein	3.92
				far upstream element (FUSE) binding prot	3.92
	424882	Al379461	Hs.153636		
	426269	H15302	Hs.168950	Homo sapiens mRNA; cDNA DKFZp566A1046 (f	3.92
	427043	AA397679	Hs.298460	ESTs	3.92
55	440404	Al015881	Hs.125616	mitochondrial ribosomal protein S5	3.92
	452762	AW501435	Hs.171409	v-akt murine thymoma viral oncogene homo	3.92
	453058	AW612293	Hs.288684	Homo sapiens cDNA FLJ11750 fis, done HE	3.92
				The state of the s	
	423583	AL122055	Hs.129836	KIAA1028 protein	3.92
<i>(</i> 0	408001	AA046458	Hs.95296	ESTs	3.92
60	419197	N48921	Hs.27441	KIAA1615 protein	3.91
	428695	Al355647	Hs.189999	purinergic receptor (family A group 5)	3.91
	401747				3.91
	410011	AB020641	Hs.57856	PFTAIRE protein kinase 1	3.91
65	432205	AI806583	Hs.125291	ESTS	3.91
65	447857	AA081218	Hs.58608	Homo sapiens cDNA FLJ14206 fis, done NT	3.91
	446494	AA463276	Hs.288906	WW Domain-Containing Gene	3.91
	409928	AL137163	Hs.57549	hypothetical protein dJ473B4	3.90
	411598	BE336654	Hs.70937	H3 histone family, member A	3.90
	424790	AL119344	Hs.13326	ESTs, Weakly similar to 2004399A chromos	3.90
	7641 <i>0</i> 0	AL113011	. 10.10020	COTO, TTOKING MINIMA IN LOUTOSON WHOMING	0.00

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	425707	AF115402	Hs.11713	E74-like factor 5 (ets domain transcript	3.90
	431325	AW026751	Hs.5794	ESTs, Weakly similar to 2109260A B cell	3.89
	451806	NM 003729	Hs.27076	RNA 3'-terminal phosphate cyclase	3.89
		1111_000120	11327010	There is thinks phosphato of Gaso	3.89
5	401045	A14/00/4700	Un 04464	thrombosopois 4	3.89
J	433023	AW864793	Hs.34161	thrombospondin 1	
	452160	BE378541	Hs.279815	cysteine sulfinic acid decarboxylase-rel	3.89
	437372	AA323968	Hs.283631	hypothetical protein DKFZp547G183	3.89
	417067	AJ001417	Hs.81086	solute carrier family 22 (extraneuronal	3.88
	410467	AF102546	Hs.63931	dachshund (Drosophila) homolog	3.88
10	422660	AW297582	Hs.237062	hypothetical protein FLJ22548 similar to	3.88
	431930	AB035301	Hs.272211	cadherin 7, type 2	3.88
	453047	AW023798	Hs.286025	ESTs	3.88
	433891	AA613792	113.200020	gb:no97h03.s1 NCI_CGAP_Pr2 Homo sapiens	3.88
		MM013/32		gualities/fluo.s1 NOI_OGAT _F12 Hollio sapiells	3.88
15	401785		11- 400004	FOT-	
15	431088	AA491824	Hs.196881	ESTs	3.88
	451952	AL120173	Hs.301663	ESTs	3.87
	422089	AA523172	Hs.103135	ESTs, Weakly similar to SFR4_HUMAN SPLIC	3.87
	452277	AL049013	Hs.28783	KIAA1223 protein	3.87
	438279	AA805166	Hs.165165	HIV-1 rev binding protein 2	3.86
20	458229	Al929602	Hs.177	phosphatidylinositol glycan, class H	3.86
	406414			,,	3.86
	417193	Al922189	Hs.288390	hypothetical protein FLJ22795	3.85
	413174	AA723564	Hs.191343	ESTs	3.85
				Homo sapiens clone TCCCTA00151 mRNA sequ	3.85
25	433332	Al367347	Hs.127809		
25	411089	AA456454	Hs.118637	cell division cycle 2-like 1 (PITSLRE pr	3.85
	412494	AL133900	Hs.792	ADP-ribosylation factor domain protein 1	3.84
	413530	AA130158	Hs.19977	ESTs, Moderately similar to ALU8_HUMAN A	3.84
	459592	AL037421	Hs.208746	ESTs, Moderately similar to pot. ORF I	3.84
	418329	AW247430	Hs.84152	cystathionine-beta-synthase	3.83
30	451468	AW503398	Hs.210047	ESTs, Moderately similar to I38022 hypot	3.83
•	434804	AA649530		gb:ns44f05.s1 NCI_CGAP_Alv1 Homo sapiens	3.83
	401819			3	3.82
	424179	F30712		Homo sapiens, clone IMAGE:4285740, mRNA	3.82
			Hs.153498	chromosome 18 open reading frame 1	3.82
35	424850	AA151057		•	3.82
22	426472	BE246138	Hs.30853	ESTs	
	426625	T78300	Hs.171409	serologically defined colon cancer antig	3.82
	427585	D31152	Hs.179729	collagen, type X, alpha 1 (Schmid metaph	3.82
	427756	Al376540	Hs.15574	ESTs ·	3.82
	444701	Al916512	Hs.198394	ESTs	3.82
40	423052	M28214	Hs.123072	RAB3B, member RAS oncogene family	3.82
	429259	AA420450	Hs.292911	ESTs, Highly similar to \$60712 band-6-pr	3.82
	416111	AA033813	Hs.79018	chromatin assembly factor 1, subunit A (3.82
	433586	T85301		gb:yd78d06.s1 Soares fetal liver spleen	3.81
	438527	Al969251	Hs.143237	RAB7, member RAS oncogene family-like 1	3.81
45	410297	AA148710	Hs.159441	lumican	3.81
73		AW117322		ESTs	3.81
	429898		Hs.42366	the state of the s	3.80
	409079	W87707	Hs.82065	interteukin 6 signal transducer (gp130,	
	419423	D26488	Hs.90315	KIAA0007 protein	3.80
~^	429643	AA455889	Hs.187548	FYVE-finger-containing Rab5 effector pro	3.80
50	431499	NM_001514	Hs.258561	general transcription factor IIB	3.80
	445060	AA830811	Hs.88808	ESTs	3.80
	449419	R34910	Hs.119172	ESTs	3.80
•	450584	AA040403	Hs.60371	ESTs -	3.80
	426137	AL040683	Hs.167031	DKFZP566D133 protein	3.79
55	420185	AL044056	Hs.158047	ESTs	3.79
55	410076	T05387	Hs.7991	ESTs	3.78
				U5 snRNP-specific 40 kDa protein (hPrp8-	3.78
	444078	BE246919	Hs.10290		
	417318	AW953937	Hs.12891	ESTs	3.78
C O	414664	AA587775	Hs.66295	multi-PDZ-domain-containing protein	3.78
60	410275	U85658	Hs.61796	transcription factor AP-2 gamma (activat	3.77
	410503	AW975746	Hs.188662	KIAA1702 protein	3.77
	434170	AA626509	Hs.122329	ESTs	3.77
	421838	AW881089	Hs.108806	Homo sapiens mRNA; cDNA DKFZp566M0947 (f	3.77
	425268	AI807883	Hs.156932	Homo sapiens cDNA FLJ20653 fis, clone KA	3.76
65	431696	AA259068	Hs.267819	protein phosphatase 1, regulatory (inhib	3.76
Ų.J	411990		Hs.31707	ESTs, Weakly similar to YEW4_YEAST HYPOT	3.76
		AW963624		CGI-49 protein	3.76
	430291	AV660345	Hs.238126		
	448779	BE042877	Hs.177135	ESTs	3.76
	452682	AA456193	Hs.155606	progesterone membrane binding protein	3.75

	452598	Al831594	Hs.68647	ESTs, Weakly similar to ALU7_HUMAN ALU S	3.75
	439498	AA908731	Hs.58297	CLLL8 protein	3.75
					3.74
	440258	Al741633	Hs.125350	ESTs	
_	456848	AL121087	Hs.296406	KIAA0685 gene product	3.74
5	415082	AA160000	Hs.137396	ESTs, Weakly similar to JC5238 galactosy	3.74
	420653	Al224532	Hs.88550	ESTs	3.74
	431637	Al879330	Hs.265960	hypothetical protein FLJ10563	3.74
	440411	N30256	Hs.156971	hypothetical protein DKFZp434G1415	3.74
	405917				3.74
10	419440	AB020689	Hs.90419	KIAA0882 protein	3.74
	451230	BE546208	Hs.26090	hypothetical protein FLJ20272	3.73
	429597	NM_003816	Hs.2442	a disintegrin and metalloproteinase doma	3.73
	430144	A1732722	Hs.187694	ERGL protein; ERGIC-53-like protein	3.72
	438394	BE379623	Hs.27693	peptidylprolyl isomerase (cyclophilin)-l	3.72
15	440527	AV657117	Hs.184164	ESTs. Moderately similar to S65657 alpha	3.72
15					
	449433	Al672096	Hs.9012	ESTs, Wealdy similar to S26650 DNA-bindi	3.72
	456228	BE503227	Hs.134759	ESTs	3.72
	448663	BE614599	Hs.106823	hypothetical protein MGC14797	3.72
	415075	L27479	Hs.77889	Friedreich ataxia region gene X123	3.72
20		Al793211	Hs.165372	ESTs, Moderately similar to ALU1_HUMAN A	3.71
20	433544				
	418293	Al224483	Hs.16063	hypothetical protein FLJ21877	3.71
	449897	AW819642	Hs.24135	transmembrane protein vezatin; hypotheti	3.71
	420297	Al628272	Hs.88323	ESTs, Weakly similar to ALU1_HUMAN ALU S	3.70
	423065	R96158	Hs.194606	Homo sapiens, clone MGC:5406, mRNA, comp	3.70
25					
23	429340	N35938	Hs.199429	Homo saplens mRNA; cDNA DKFZp434M2216 (f	3.70
	437777	AA768098	Hs.189079	ESTs	3.70
	440351	AF030933	Hs.7179	RAD1 (S. pombe) homolog	3.70
	443603	BE502601	Hs.134289	ESTs, Weakly similar to KIAA1063 protein	3.70
	446965	BE242873	Hs.16677	WD repeat domain 15	3.70
20					
30	412350	A1659306	Hs.73826	protein tyrosine phosphatase, non-recept	3.70
	433852	Al378329	Hs.126629	ESTs	3.70
	433142	AL120697	Hs.110640	ESTs	3.69
	419994	AA282881	Hs.190057	ESTs .	3.69
	412628	Al972402	Hs.173902	hypothetical protein MGC2648	3.69
35					
22	431416	AA532718	Hs.178604	ESTs	3.69
	439444	Al277652	Hs.54578	ESTs, Weakly similar to I38022 hypotheti	3.68
	414709	AA704703	Hs.77031	Sp2 transcription factor	3.68
	447397	BE247676	Hs.18442	E-1 enzyme	3.68
	405718				3.68
40		ALIOTECOE	Un 155174	CDCE (call distains guals E. C. nombo h	3.68
40	425217	AU076696	Hs.155174	CDC5 (cell division cycle 5, S. pombe, h	
	442242	AV647908	Hs.90424	Homo sapiens cDNA: FLJ23285 fis, clone H	3.68
	424690	BE538356	Hs.151777	eukaryotic translation initiation factor	3.68
	421734	Al318624	Hs.107444	Homo sapiens cDNA FLJ20562 fis, clone KA	3.67
	427221	L15409	Hs.174007	von Hippel-Lindau syndrome	3.67
45					
43	439864	AI720078	Hs.291997	ESTs, Weakly similar to A47582 B-cell gr	3.66
	402408			·	3.66
	426327	W03242	Hs.44898	Homo sapiens done TCCCTA00151 mRNA sequ	3.66
	427119	AW880562	Hs.114574	ESTs	3.66
	427356	AW023482	Hs.97849	ESTs	3.66
50					
20	452946	X95425	Hs.31092	EphA5	3.66
	419078	M93119	Hs.89584	Insulinoma-associated 1	3.66
			11- 400000	FOY-	3.65
	416295	A1064824	MS.193385	ESTs	0.00
	416295 427144		Hs.193385 Hs.2126		
	427144	X95097	Hs.2126	vasoactive intestinal peptide receptor 2	3.65
<i>==</i>	427144 447500	X95097 Al381900	Hs.2126 Hs.159212	vasoactive intestinal peptide receptor 2 ESTs	3.65 3.65
55	427144 447500 453127	X95097 Al381900 Al696671	Hs.2126 Hs.159212 Hs.294110	vasoactive intestinal peptide receptor 2 ESTs ESTs	3.65 3.65 3.65
55	427144 447500	X95097 Al381900	Hs.2126 Hs.159212	vasoactive intestinal peptide receptor 2 ESTs	3.65 3.65
55	427144 447500 453127 423396	X95097 Al381900 Al696671	Hs.2126 Hs.159212 Hs.294110	vasoactive intestinal peptide receptor 2 ESTs ESTs	3.65 3.65 3.65
55	427144 447500 453127 423396 419346	X95097 Al381900 Al696671 Al382555 Al830417	Hs.2126 Hs.159212 Hs.294110 Hs.127950	vasoactive intestinal peptide receptor 2 ESTs ESTs bromodomain-containing 1 polybromo 1	3.65 3.65 3.65 3.65 3.64
55	427144 447500 453127 423396 419346 441540	X95097 Al381900 Al696671 Al382555 Al830417 C01367	Hs.2126 Hs.159212 Hs.294110 Hs.127950	vasoactive intestinal peptide receptor 2 ESTs ESTs bromodomain-containing 1 polybromo 1 ESTs	3.65 3.65 3.65 3.65 3.64 3.64
	427144 447500 453127 423396 419346 441540 446501	X95097 Al381900 Al696671 Al382555 Al830417 C01367 Al302616	Hs.2126 Hs.159212 Hs.294110 Hs.127950 Hs.127128 Hs.150819	vasoactive intestinal peptide receptor 2 ESTs ESTs bromodomain-containing 1 polybromo 1 ESTs ESTs	3.65 3.65 3.65 3.65 3.64 3.64
55 60	427144 447500 453127 423396 419346 441540 446501 459527	X95097 Al381900 Al696671 Al382555 Al830417 C01367 Al302616 AW977556	Hs.2126 Hs.159212 Hs.294110 Hs.127950 Hs.127128 Hs.150819 Hs.291735	vasoactive intestinal peptide receptor 2 ESTs ESTs bromodomain-containing 1 polybromo 1 ESTs ESTs ESTs ESTs ESTs ESTs, Weakty similar to 178885 serine/th	3.65 3.65 3.65 3.65 3.64 3.64 3.64 3.63
	427144 447500 453127 423396 419346 441540 446501	X95097 Al381900 Al696671 Al382555 Al830417 C01367 Al302616	Hs.2126 Hs.159212 Hs.294110 Hs.127950 Hs.127128 Hs.150819	vasoactive intestinal peptide receptor 2 ESTs ESTs bromodomain-containing 1 polybromo 1 ESTs ESTs ESTs ESTs ESTs ESTs, Weakty similar to 178885 serine/th	3.65 3.65 3.65 3.65 3.64 3.64
	427144 447500 453127 423396 419346 441540 446501 459527 446320	X95097 Al381900 Al696671 Al382555 Al830417 C01367 Al302616 AW977556 AF126245	Hs.2126 Hs.159212 Hs.294110 Hs.127950 Hs.127128 Hs.150819 Hs.291735 Hs.14791	vasoactive intestinal peptide receptor 2 ESTs ESTs bromodomain-containing 1 polybromo 1 ESTs ESTs ESTs ESTs ESTs, Weakly similar to I78885 serine/th acyl-Coenzyme A dehydrogenase family, me	3.65 3.65 3.65 3.65 3.64 3.64 3.63 3.63
	427144 447500 453127 423396 419346 441540 446501 459527 446320 435706	X95097 Al381900 Al696671 Al382555 Al830417 C01367 Al302616 AW977556	Hs.2126 Hs.159212 Hs.294110 Hs.127950 Hs.127128 Hs.150819 Hs.291735	vasoactive intestinal peptide receptor 2 ESTs ESTs bromodomain-containing 1 polybromo 1 ESTs ESTs ESTs ESTs ESTs ESTs, Weakty similar to 178885 serine/th	3.65 3.65 3.65 3.64 3.64 3.63 3.63 3.63
	427144 447500 453127 423396 419346 441540 446501 459527 446320 435706 400110	X95097 Al381900 Al696671 Al382555 Al830417 C01367 Al302616 AW977556 AF126245 W31254	Hs.2126 Hs.159212 Hs.294110 Hs.127950 Hs.127128 Hs.150819 Hs.291735 Hs.14791 Hs.7045	vasoactive intestinal peptide receptor 2 ESTs ESTs bromodomain-containing 1 polybromo 1 ESTs ESTs ESTs ESTs ESTs ESTs, Weakly similar to I78885 serine/th acyl-Coenzyme A dehydrogenase family, me GL004 protein	3.65 3.65 3.65 3.64 3.64 3.63 3.63 3.63 3.63
60	427144 447500 453127 423396 419346 441540 446501 459527 446320 435706 400110 410313	X95097 Al381900 Al696671 Al382555 Al830417 C01367 Al302616 AW977556 AF126245 W31254	Hs.2126 Hs.159212 Hs.294110 Hs.127950 Hs.127128 Hs.150819 Hs.291735 Hs.14791 Hs.7045 Hs.185683	vasoactive intestinal peptide receptor 2 ESTs ESTs bromodomain-containing 1 polybromo 1 ESTs ESTs ESTs, Weakly similar to I78885 serine/th acyi-Coenzyme A dehydrogenase family, me GL004 protein	3.65 3.65 3.65 3.64 3.64 3.63 3.63 3.63 3.63 3.62
	427144 447500 453127 423396 419346 441540 446501 459527 446320 435706 400110	X95097 Al381900 Al696671 Al382555 Al830417 C01367 Al302616 AW977556 AF126245 W31254 R10305 BE465243	Hs.2126 Hs.159212 Hs.294110 Hs.127950 Hs.127128 Hs.150819 Hs.291735 Hs.14791 Hs.7045 Hs.185683 Hs.12664	vasoactive intestinal peptide receptor 2 ESTs ESTs bromodomain-containing 1 polybromo 1 ESTs ESTs ESTs, Weakly similar to I78885 serine/th acyl-Coenzyme A dehydrogenase family, me GL004 protein ESTs ESTs	3.65 3.65 3.65 3.64 3.64 3.63 3.63 3.63 3.63 3.62 3.62 3.62
60	427144 447500 453127 423396 419346 441540 446501 459527 446320 435706 400110 410313	X95097 Al381900 Al696671 Al382555 Al830417 C01367 Al302616 AW977556 AF126245 W31254	Hs.2126 Hs.159212 Hs.294110 Hs.127950 Hs.127128 Hs.150819 Hs.291735 Hs.14791 Hs.7045 Hs.185683	vasoactive intestinal peptide receptor 2 ESTs ESTs bromodomain-containing 1 polybromo 1 ESTs ESTs ESTs, Weakly similar to I78885 serine/th acyl-Coenzyme A dehydrogenase family, me GL004 protein ESTs ESTs	3.65 3.65 3.65 3.64 3.64 3.63 3.63 3.63 3.63 3.62
60	427144 447500 453127 423396 419346 441540 446501 459527 446320 435706 400110 410313 414713 436279	X95097 AI381900 AI696671 AI382555 AI830417 C01387 AI302616 AW977556 AF126245 W31254 R10305 BE465243 AW900372	Hs.2126 Hs.159212 Hs.294110 Hs.127950 Hs.127128 Hs.150819 Hs.291735 Hs.14791 Hs.7045 Hs.185683 Hs.12664 Hs.180793	vasoactive intestinal peptide receptor 2 ESTs ESTs bromodomain-containing 1 polybromo 1 ESTs ESTs ESTs, Weakly similar to I78885 serine/th acyl-Coenzyme A dehydrogenase family, me GL004 protein ESTs ESTs ESTs, Weakly similar to S65657 elpha-1C-	3.65 3.65 3.65 3.64 3.64 3.63 3.63 3.63 3.62 3.62 3.62 3.62
60	427144 447500 453127 423396 419346 441540 446501 459527 446320 435706 400110 410313 414713 436279 439818	X95097 AI381900 AI696671 AI382555 AI830417 C01367 AI302616 AW977556 AF126245 W31254 R10305 BE465243 AW900372 AL360137	Hs.2126 Hs.159212 Hs.294110 Hs.127950 Hs.127128 Hs.150819 Hs.291735 Hs.14791 Hs.7045 Hs.185683 Hs.12664 Hs.180793 Hs.19934	vasoactive intestinal peptide receptor 2 ESTs ESTs bromodomain-containing 1 polybromo 1 ESTs ESTs ESTs, Weakly similar to I78885 serine/th acyl-Coenzyme A dehydrogenase family, me GL004 protein ESTs ESTs ESTs ESTs ESTs ESTs Weakly similar to S65657 alpha-1C- Homo sapiens mRNA full length insert cDN	3.65 3.65 3.65 3.64 3.64 3.63 3.63 3.63 3.62 3.62 3.62 3.62
60	427144 447500 453127 423396 419346 441540 446501 459527 446320 435706 400110 410313 414713 436279	X95097 AI381900 AI696671 AI382555 AI830417 C01387 AI302616 AW977556 AF126245 W31254 R10305 BE465243 AW900372	Hs.2126 Hs.159212 Hs.294110 Hs.127950 Hs.127128 Hs.150819 Hs.291735 Hs.14791 Hs.7045 Hs.185683 Hs.12664 Hs.180793	vasoactive intestinal peptide receptor 2 ESTs ESTs bromodomain-containing 1 polybromo 1 ESTs ESTs ESTs, Weakly similar to I78885 serine/th acyl-Coenzyme A dehydrogenase family, me GL004 protein ESTs ESTs ESTs, Weakly similar to S65657 elpha-1C-	3.65 3.65 3.65 3.64 3.64 3.63 3.63 3.63 3.62 3.62 3.62 3.62

	434194	AF119847	Hs.283940	Homo sapiens PRO1550 mRNA, partial cds	3.62
	404939				3.62
	408101	AW968504	Hs.123073	CDC2-related protein kinase 7	3.62
_	435846	AA700870	Hs.14304	ESTs	3.61
5	432833	N51075	Hs.47191	ESTs	3.61
	427276	AA400269	Hs.49598	ESTs	3.61
	433495	AW373784	Hs.71	alpha-2-glycoprotein 1, zinc	3.60
	403137				3.60
10	404165				3.60
5 10	409571	AA504249	Hs.187585	ESTs	3.60
	410561	BE540255	Hs.6994	Homo sapiens cDNA: FLi22044 fis, clone H	3.60
	412924	BE018422	Hs.75258	H2A histone family, member Y	3.60
	434228	Z42047	Hs.283978	Homo sapiens PRO2751 mRNA, complete cds	3.60
15	436797	AA731491	Hs.178518	hypothetical protein MGC14879	3.60
13	437162	. AW005505	Hs.5464	thyroid hormone receptor coactivating pr	3.60
	437444	H46008	Hs.31518	ESTs	3.60
	404210	BE270828	Hs.131740	Home conjuga eDNA: EL MOSSO fin elega U	3.59 3.59
	446157 437587	AI591222	Hs.122421	Homo sapiens cDNA: FLJ22562 fis, clone H Human DNA sequence from clone RP1-187J11	3.58
20	43/36/	AA987927	Hs.131740	Homo sapiens cDNA: FLJ22562 fis, done H	3.57
20	452226	AA024898	Hs.296002	ESTs	3.56
	452220	AF291664	Hs.204732	matrix metalloproteinase 26	3.56
	452501	AB037791	Hs.29716	hypothetical protein FLJ 10980	3.56
		AA830050	Hs.124344	ESTs	3.56
25	428647 422443	NM_014707	Hs.116753	histone deacetylase 78	3.55
23	447966	AA340605	Hs.105887	ESTs, Weakly similar to Homolog of rat Z	3.55
	420892	AW975076	Hs.172589	nuclear phosphoprotein similar to S. cer	3.55
	420230	AL034344	Hs.298020	forkhead box C1	3.55
	418428	Y12490	Hs.85092	thyroid hormone receptor interactor 11	3.54
30	428949	AA442153	Hs.104744	hypothetical protein DKFZp434J0617	3.54
-	444929	Al685841	Hs.161354	ESTs	3.54
	433339	AF019226	Hs.8036	giloblastoma overexpressed	3.54
	424369	R87622	Hs.26714	KIAA1831 protein	3.54
	433002	AF048730	Hs.279906	cyclin T1	3.53
35	435425	H16263	Hs.31416	ESTs	3.53
	415621	Al648602	Hs.131189	ESTs	3.53
	416974	AF010233	Hs.80667	RALBP1 associated Eps domain containing	3.53
	405793			,	3.52
	409770	AW499536		gb:UI-HF-BR0p-aff-c-12-0-UI.r1 NIH_MGC_5	3.52
40	425305	AA363025	Hs.155572	Human clone 23801 mRNA sequence	3.52
	428939	AW236550	Hs.131914	ESTs	3.52
	438388	AA806349	Hs.44698	ESTs	3.52
	443703	AV646177	Hs.213021	ESTs	3.52
	457940	AL360159	Hs.30445	Homo sapiens TRIpartite motif protein ps	3.52
45	402444				3.52
	409643	AW450866	Hs.257359	ESTs	3.51
	418250	U29926	Hs.83918	adenosine monophosphate deaminase (isofo	3.51
•	432745	Al821926	Hs.269507	gb:nt78f05.x5 NCI_CGAP_Pr3 Homo sapiens	3.51
50	414222	AL135173	Hs.878	sorbitol dehydrogenase	3.51
50	430061	AB037817	Hs.230188	KIAA1396 protein	3.51
	421491	H99999	Hs.42736	ESTs	3.50
	422384	AA224077	Hs.42438	Sm protein F	3.50
	434565	T52172	H- 474004	ESTs -	3.50
55	438379	N23018	Hs.171391	C-terminal binding protein 2	3.50
55	439741	BE379646 R37010	Hs.6904	Homo sapiens mRNA full length insert cDN Homo sapiens cDNA: FLJ22806 fis, clone K	3.50 3.50
	447311 447805	AW627932	Hs.33417 Hs.19614	gemin4	3.50
	454265	H03556	Hs.300949	ESTs, Weakly similar to thyrold hormone	3.50
	418838	AW385224	Hs.35198	ectonucleotide pyrophosphatase/phosphodi	3.50
60	448804	AW512213	Hs.42500	ADP-ribosylation factor-like 5	3.50
	409617	BE003760	Hs.55209	Homo sapiens mRNA; cDNA DKFZp434K0514 (f	3.49
	434075	AW003416	Hs.160604	ESTs	3.49
	444190	AI878918	Hs.10526	cysteine and glycine-rich protein 2	3.49
	435017	AA336522	Hs.12854	angiotensin II, type I receptor-associat	3.48
65	423445	NM_014324	Hs.128749	alpha-methylacyl-CoA racemase	3.48
	420271	AI954365	Hs.42892	ESTs	3.48
	443684	AI681307	Hs.166674	ESTs	3.48
	444168	AW379879		gb:RC1-HT0256-081199-011-f01 HT0256 Homo	3.48
	446074	AA079799	Hs.29263	hypothetical protein FLJ11896	3.48
			• •	••	

5	452582 431542 432697 435572 407192 413435 447210	AL137407 H63010 AW975050 AW975339 AA609200 X51405 AF035269	Hs.29911 Hs.5740 Hs.293892 Hs.239828 Hs.75360 Hs.17752	Homo sapiens mRNA; cDNA DKFZp434M232 (fr ESTs ESTs, Weakly similar to ALU4_HUMAN ALU S ESTs, Weakly similar to GAG2_HUMAN RETRO gb:af12e02.s1 Soares_testis_NHT Homo sap carboxypeptidase E phosphatidytserine-specific phospholipas	3.48 3.48 3.47 3.47 3.46 3.46
	447958	AW796524	Hs.68644	Homo sapiens microsomal signal peptidase	3.46
10	425312	AA354940	Hs.145958	ESTs	3.46 3.46
IU	442007	AA301116	Hs.142838	nucleolar phosphoprotein Nopp34	3.45
	417455	AW007066	Hs.18949	ESTs, Weakly similar to CA2B_HUMAN COLLA	
	426931	NM_003416	Hs.2076	zinc finger protein 7 (KOX 4, clone HF.1	3.45
	408739	W01556	Hs.238797	ESTs, Moderately similar to I38022 hypot	3.45
	436024	Ai800041	Hs.190555	ESTs	3.45
15	408418	AW963897	Hs.44743	KIAA1435 protein	3.45
	409151	AA306105	Hs.50785	SEC22, vesicle trafficking protein (S. c	3.44
	418626	AW299508	Hs.135230	ESTs	3.44
	420560	AW207748	Hs.59115	ESTs	3.44
	420686	A1950339	Hs.40782	ESTs .	3.44
20	428870	AA436831	Hs.36049	ESTs	3.44
	436754	Al061288	Hs.133437	ESTs	3.44
	437960	A1669586	Hs.222194	ESTs	3.44
	452300	AW628045	Hs.28896	Homo sapiens mRNA full length insert cDN	3.44
	421887	AW161450	Hs.109201	CGI-86 protein	3.44
25				• • • • • • • • • • • • • • • • • •	

TABLE 5A shows the accession numbers for those primekeys lacking a unigeneID in Tables 5, 6, and 7. For each probeset we have listed the gene cluster number from which the oligonucleotides were designed. Gene clusters were compiled using sequences derived from Genbank ESTs and mRNAs. These sequences were clustered based on sequence similarity using Clustering and Alignment Tools (DoubleTwist, Oakland California). The Genbank accession numbers for sequences comprising each cluster are listed in the "Accession" column.

5

	Pkey:		Unique Eos probeset identifier number
	CAT number:		Gene cluster number
	Accession:		Genbank accession numbers
10	Pkey	CAT number	Accession
	407596	1003489_1	R86913 R86901 H25352 R01370 H43764 AW044451 W21298
٠,,	408432	1058667_1	AW195262 R27868 AW811262
15	409752	115301_1	AW963990 AA078196 AW749482 AA077468 BE151571 AA376917
	409770	1154048_1	AW499536 AW499553 AW502138 AW499537 AW502136 AW501743
	411440	124577_1	AW749402 AW749403 Z45743 R80376 AA093358
	411479	1247077_1	AW848047 AW848202 AW848631 AW848142 AW848702 AW848121 AW848632 AW848140 AW848571 AW848009 AW848067 AW848069 AW848905 AW848214
20	411624	1252166_1	BE145964 BE146286 AW854564
	412991	134248_1	AW949013 AA126111
	414269	143133_1	AA298489 AA137165
	415123	1523390_1	D60925 D60828 D80787
	415715	1548818_1	F30364 F36559 T15435
25	416288	1585983_1	H51299 H44619 H46391 R86024 H51892 T72744
	416289	1586037_1	W26333 R05358 H44682
	417730	1695795_1	Z44761 R25801 R11926 R35604
	418636	177402_1	AW749855 AA225995 AW750208 AW750206
••	419346	184129_1	Al830417 AA236612
30	419536	185688_1	AA603305 AA244095 AA244183
	420111	190755_1	AA255652 AA280911 AW967920 AA262684
	422219	213547_1	AW978073 AW978072 AA807550 AA306567
	424179	236389_1	F30712 F35665 AW263888 Al904014 Al904018 AA336927 AA336502
25	424242	237181_1	AA337476 AW966227 AA450376 AW960222 AA381051
35	428002	285602_1	AA418703 AA418711 BE071915 BE071920 BE071912
	429163	300543_1	AA884766 AW974271 AA592975 AA447312
	432189	342819_1	AA527941 AI810608 AI620190 AA635266
	432340	345248_1 345469_1	AA534222 AA632632 T81234 AA534490 AM977040 AM9770323
40	432363 432966	356839_1	AA534489 AW970240 AW970323 AA650114 AW974148 AA572946
40	432900 433586	370470 1	T85301 AW517087 AA601054 BE073959
	433641	37186_1	AF080229 AF080231 AF080230 AF080232 AF080233 AF080234 BE550633 Al636743 AW614951 BE467547
	433041	3/100_1	A1680833 A1633818 N29986 U87592 U87593 U87590 U87591 S46404 U87587 AA463992 AW206802 A1970376
			AI583718 AI672574 N25695 AW665466 AI618326 AA126128 AI480345 AW013827 AA248638 AI214968
45			AA204735 AA207155 AA206262 AA204833 AW003247 AW496808 Al080480 Al631703 Al651023 Al867418
••			AW818140 AA502500 Al206199 Al671282 Al352545 BE501030 Al652535 BE465762 AA206331 AW451866
			AA471088 AA206342 AA204834 AA206100 AW021661 AA332922 N66048 AA703396 H92278 AW139734
			H92683 U87589 U87595 H69001 U87594 BE466420 Al624817 BE466611 Al206344 AA574397 AA348354
			Al493192
50	433687	373061_1	AA743991 AA604852 AW272737
	433891	376239 1	AA613792 AW182329 T05304 AW858385
	434415	385931_1	BE177494 AW276909 AA632849
	434565	38898_1	T52172 AF147324 T52248
	434804	393481_1	AA649530 AA659316 H64973
55	437113	433234_1	AA744693 AW750059
	444168	593829_1	AW379879 Al126285 H12014
	448212	755099_1	Al475858 AW969013
	448310	757918_1	Al480316 AW847535
	451746	883303_1	M86178 Al813822 D56993
		_	

	452560	922216_1	BE077084 AW139963 AW863127 AW806209 AW806204 AW806205 AW806206 AW806211 AW806212
			AW806207 AW806208 AW806210 Al907497
	452712	928309_1	AW838616 AW838660 BE144343 Al914520 AW888910 BE184854 BE184784
	453773	980699_1	AL133761 AL133767
5	455276	1272541_1	BE176479 BE176678 BE176357 BE176550 AW886079 BE176676 BE176615 BE176555 BE176489 BE176610
			BE176362
	455309	1278153_1	AW894017 AW893956 AW894032

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7711566

TABLE 5B shows the genomic positioning for those primekeys lacking unigene ID's and accession numbers in Tables 5, 6, and 7. For each predicted exon, we have listed the genomic sequence source used for prediction. Nucleotide locations of each predicted exon are also listed.

10	Pkey: Ref:	Unique number corresponding to an Eos probeset Sequence source. The 7 digit numbers in this column are Genbank Identifier (GI) numbers. "Dunham I. et al." refers to the publication entitled "The DNA sequence of human chromosome 22." Dunham I. et al., Nature (1999) 402:489-495.
	Strand:	Indicates DNA strand from which exons were predicted.
	Nt_position:	Indicates nucleotide positions of predicted exons.

15				
	Pkey	Ref	Strand	Nt_position
	401045	8117619	Plus	90044-90184,91111-91345
20	401424	8176894	Plus	24223-24428
	401451	6634068	Minus	119926-121272
	401714	6715702	Plus	96484-96681
	401747	9789672	Minus	118596-118816,119119-119244,119609-119761,120422-120990,130161-130381,130468-130593,131097-
				131258,131866-131932,132451-132575,133580-134011
25	401785	7249190	Minus	165776-165996,166189-166314,166408-166569,167112-167268,167387-167469,168634-168942
	401819	7467933	Minus	28217-28486
	402408	9796239	Minus	110326-110491
	402444	9796614	Plus	28391-28517
	402791	6137008	Minus	51036-51207
30	403047	3540153	· Minus	59793-59968
	403137	9211494	Minus	92349-92572,92958-93084,93579-93712,93949-94072,94591-94748,95214-95337
	403721	7528046	Minus	156647-157366
	403764	7717105	Minus	118692-118853
	403797	8099896	Minus	123065-125008
35	404165	9926489	Minus	69025-69128
	404210	5006246	Plus	169926-170121
	404253	9367202	Minus	55675-56055
	404561	9795980	Minus	69039-70100
	404571	7249169	Minus	112450-112648
40	404721	9856648	Minus	173763-174294
	404915	7341766	Minus	100915-101087
	404939	6862697	Plus	175318-175476
	405403	6850244	Minus	37491-37670,40951-41031
	405685	4508129	Minus	37956-38097
45	405718	9795467	Plus	113080-113266
	405793	1405887	Minus	89197-89453
	405876	6758747	Plus	39694-40031
	405917	7712162	Minus	106829-107213
	406414	9256407	Plus	49593-49850

106956-107121

TABLE 6:286 GENES ENCODING EXTRACELLULAR OR CELL SURFACE PROTEINS UP-REGULATED IN PROSTATE CANCER COMPARED TO NORMAL ADULT TISSUES

Table 6 shows 286 genes up-regulated in prostate cancer compared to normal adult tissues that are likely to be extracellular or cell-surface proteins. These were selected as for Table 5 and the predicted protein contained a structural domain that is indicative of extracellular localization (e.g. egf, 7tm domains).

				•					
	Pkey:		Unique Eos pa	robeset identifier number					
	ExAccn:			ession number, Genbank accession number					
	UnigenelD								
	•			Unigene number Unigene gene title					
	Unigene Ti	ue.							
	R1:		Hatio of tumor	to normal tissue					
10	Pkey	ExAcon	UnigenelD	Uningene Title	R1				
	409361	NM_005982	Hs.54416	sine oculis homeobox (Drosophila) homolo	48.28				
	409731	AA125985	Hs.56145	thymosin, beta, identified in neuroblast	45.24				
	400298	AA032279	Hs.61635	six transmembrane epithelial antigen of	43.48				
15	420154	Al093155	Hs.95420	JM27 protein	41.12				
13									
	426747	AA535210	Hs.171995	kallikrein 3, (prostate specific antigen	31.80				
	400299	X07730	Hs.171995	kallikrein 3, (prostate specific antigen	24.91				
	425075	AA506324	Hs.1852	acid phosphatase, prostate	24.23				
••	424846	AU077324	Hs.1832	neuropeptide Y	23.57				
20	405685				20.90				
	420757	X78592	Hs.99915	androgen receptor (dihydrotestosterone r	19.72				
	418994	AA296520	Hs.89546	selectin E (endothelial adhesion molecul	19.56				
	452792	AB037765	Hs.30652	KIAA1344 protein	17.39				
	445472	AB006631	Hs.12784	Homo sapiens mRNA for KIAA0293 gene, par	17.00				
25	414565	AA502972	Hs.183390	hypothetical protein FLJ13590	16.82				
	431716	D89053	Hs.268012						
				fatty-acid-Coenzyme A ligase, long-chain	16.60				
	408430	S79876	Hs.44926	dipeptidylpeptidase IV (CD26, adenosine	16.28				
	408000	L11690	Hs.620	bullous pemphigoid antigen 1 (230/240kD)	15.54				
20	430226	BE245562	Hs.2551	adrenergic, beta-2-, receptor, surface	15.40				
30	444484	AK002126	Hs.11260	hypothetical protein FLJ11264	14.76				
	418601	AA279490	Hs.86368	calmegin	14.56				
	448999	AF179274	Hs.22791	transmembrane protein with EGF-like and	14.55				
	416182	NM_004354	Hs.79069	cyclin G2	12.94				
	420544	AA677577	Hs.98732	Homo sapiens Chromosome 16 BAC clone CIT	12.79				
35	445413	AA151342	Hs.12677	CGI-147 protein	12.64				
	453930	AA419466	Hs.36727	hypothetical protein FLJ10903	12.22				
	440286	U29589	Hs.7138	cholinergic receptor, muscarinic 3	12.04				
	452784	BE463857	Hs.151258	hypothetical protein FLJ21062	11.86				
	450203	AF097994	Hs.301528						
40				L-kynurenine/alpha-aminoadipate aminotra	11.68				
40	448045	AJ297436	Hs.20166	prostate stem cell antigen	11.51				
	449650	AF055575	Hs.23838	calcium channel, voltage-dependent, L ty	11.18				
	420381	D50640	Hs.337616	phosphodiesterase 3B, cGMP-inhibited	11.10				
	425665	AK001050	Hs.159066	hypothetical protein FLJ10188	11.08				
	425710	AF030880	Hs.159275	solute carrier family, member 4	11.08				
45	428728	NM_016625	Hs.191381	hypothetical protein	11.04				
	407021	U52077		gb:Human mariner1 transposase gene, comp	11.02				
	410733	D84284	Hs.66052	CD38 antigen (p45)	11.02				
	452340	NM 002202	Hs.505	ISL1 transcription factor, LIM/homeodoma	10.85				
	428819	AL135623	Hs.193914	KIAA0575 gene product	10.48				
50	421991	NM_014918	Hs.110488	KIAA0990 protein	10.04				
50	431217	NM_013427							
		R27496	Hs.250830	Rho GTPase activating protein 6	9.75				
	421470		Hs.1378	annexin A3	9.64				
	409262	AK000631	Hs.52256	hypothetical protein FLJ20624	9.45				
	435980	AF274571	Hs.129142	deoxyribonuclease II beta	9.24				
55	421246	AW582962	Hs.102897	CGI-47 protein	9.20				
	410001	AB041036	Hs.57771	kallikrein 11	9.03				
	441791	AW372449	Hs.175982	hypothetical protein FLJ21159	9.02				
				••					

	404574				8.66
	404571	AW967956	He 1006/0	ESTs, Weakly similar to AF108460 1 ubinu	8.56
	456497		Hs.123648	interleukin 6 (interleron, beta 2)	8.36
	419968 433172	X04430 AB037841	Hs.93913 Hs.102652	hypothetical protein ASH1	8.30
5	433172	BE218919	Hs.118793	hypothetical protein FLJ10688	8.27
5	427674	NM_003528	Hs.2178	H2B histone family, member Q	8.20
	404915	MILOOOSEO	1132170	The Holder tamey, money	8.08
	452259	AA317439	Hs.28707	signal sequence receptor, gamma (translo	8.06
	452891	N75582	Hs.212875	ESTs, Weakly similar to DYH9_HUMAN CILIA	8.02
10	439731	Al953135	Hs.45140	hypothetical protein FLJ14084	7.98
	419839	U24577	Hs.93304	phospholipase A2, group VII (platelet-ac	7.68
	420120	AL049610	Hs.95243	transcription elongation factor A (SII)-	7.64
	424099	AF071202	Hs.139336	ATP-binding cassette, sub-family C (CFTR	7.64
	448706	AW291095	Hs.21814	interleukin 20 receptor, alpha	7.52
15	410227	AB009284	Hs.61152	exostoses (multiple)-like 2	7.49
	425211	M18667	Hs.1867	progastricsin (pepsinogen C)	7.35 7.28
	441736	AW292779	Hs.169799	ESTs	7.20
	419991	AJ000098	Hs.94210 Hs.154196	eyes absent (Drosophila) homolog 1 E4F transcription factor 1	7.20
20	425018 424560	BE245277 AA158727	Hs.150555	protein predicted by clone 23733	7.18
20	424560	AA191493	Hs.48778	niban protein	7.10
	421566	NM_000399	Hs.1395	early growth response 2 (Krox-20 (Drosop	7.04
	431725	X65724	Hs.2839	Norrie disease (pseudoglioma)	6.98
	425782	U66468	Hs.159525	cell growth regulatory with EF-hand doma	6.85
25	427408	AA583206	Hs.2156	RAR-related orphan receptor A	6.79
	435604	AA625279	Hs.26892	uncharacterized bone marrow protein BM04	6.73
	415874	AF091622	Hs.78893	KIAA0244 protein	6.54
	401451				6.52
20	431778	AL080276	Hs.268562	regulator of G-protein signalling 17	6.51
30	409089	NM_014781	Hs.50421	KIAA0203 gene product	6.50 6.49
	431992	NM_002742	Hs.2891	protein kinase C, mu	6.42
	404253 404553	AF026692	Hs.105700	secreted frizzled-related protein 4	6.41
	421552 416806	NM_000288	Hs.79993	peroxisomal biogenesis factor 7	6.38
35	431958	X63629	Hs.2877	cadherin 3, type 1, P-cadherin (placenta	6.30
55	439366	AF100143	Hs.6540	fibroblast growth factor 13	6.30
	416836	D54745	Hs.80247	cholecystokinin	6.30
	433383	AF034837	Hs.192731	double-stranded RNA specific adenosine d	6.29
	450728	AW162923	Hs.25363	presenilin 2 (Alzheimer disease 4)	6.25
40	413384	NM_000401	Hs.75334	exostoses (multiple) 2	6.22
	423349	AF010258	Hs.127428	homeo box A9	6.20
	424800	AL035588	Hs.153203	MyoD family inhibitor	6.18
	425451	AF242769	Hs.157461	mesenchymal stem cell protein DSC54	6.14 6.00
45	447359	NM_012093	Hs.18268 Hs.66744	adenylate kinase 5 twist (Drosophila) homolog (acrocephalos	5.97
43	410889 408829	X91662 NM_006042	Hs.48384	heparan sulfate (glucosamine) 3-0-sulfot	5.94
	453911	AW503857	Hs.4007	Sarcolemmal-associated protein	5.94
	408875	NM_015434	Hs.48604	DKFZP434B168 protein	5.92
	450480	X82125	Hs.25040	zinc finger protein 239	5.90
50	451684	AF216751	Hs.26813	CDA14	5.88
	400301	X03635	Hs.1657	estrogen receptor 1	5.78
	415077	L41607	Hs.934	glucosaminyl (N-acetyl) transferase 2, I	5.74
	418852	BE537037	Hs.273294	hypothetical protein FLJ20069	5.72
	446867	AB007891	Hs.16349	KIAA0431 protein	5.72
55	410232	AW372451	Hs.61184	CGI-79 protein	5.70
	422762	AL031320	Hs.119976	Human DNA sequence from clone RP1-20N2 o	5.70 5.70
	450616	AL133067	Hs.302689	hypothetical protein chromosome 11 open reading frame 8	5.65
	408621	AI970672	Hs.46638 Hs.6641	kinesin family member 5C	5.64
60	439671 410196	AW162840 Al936442	Hs.59838	hypothetical protein FLJ10808	5.60
00	429170	NM_001394	Hs.2359	dual specificity phosphatase 4	5.60
	440738	A1004650	Hs.225674	WD repeat domain 9	5.60
	414342	AA742181	Hs.75912	KIAA0257 protein	5.59
	422634	NM_016010	Hs.118821	CGI-62 protein	5.56
65	400268			•	5.55
	439569	AW602166	Hs.222399	CEGP1 protein	5.51
	452823	AB012124	Hs.30696	transcription factor-like 5 (basic helix	5.48
	431938	AA938471	Hs.54431	specific granule protein (28 kDa); cyste	5.44
•	427638	AA406411	Hs.208341	ESTs, Weakly similar to KIAA0989 protein	5.42

	421264	AL039123	Hs.103042	microtubule-associated protein 1B	5.38
	421685	AF189723	Hs.106778	ATPase, Ca++ transporting, type 2C, memb	5.37
	421987	Al133161	Hs.286131	CGI-101 protein	5.3 6
	422806	BE314767	Hs.1581	glutathione S-transferase theta 2	5.34
-5	432281	AK001239	Hs.274263	hypothetical protein FLJ10377	5.32
	451982	F13036	Hs.27373	Homo sapiens mRNA; cDNA DKFZp564O1763 (f	5.32
	444042	NM_004915	Hs.10237	ATP-binding cassette, sub-family G (WHIT	5.31
	447752	M73700	Hs.105938	lactotransferrin	5.29
	451418	BE387790	Hs.26369	hypothetical protein FLJ20287	5.22
10	428593	AW207440	Hs.185973	degenerative spermatocyte (homolog Droso	5.21
	447541	AK000288	Hs.18800	hypothetical protein FLJ20281	5.18
	459294	AW977286	Hs.17428	RBP1-like protein	5.16
	424692	AA429834	Hs.151791	KIAA0092 gene product	5.15
	416434	AW163045	Hs.79334	nuclear factor, interleukin 3 regulated	5.11
15	410268	AA316181	Hs.61635	six transmembrane epithelial antigen of	5.10
	417517	AF001176	Hs.82238	POP4 (processing of precursor, S. cerev	5.10
	453616	NM_003462	Hs.33846	dynein, axonemal, light intermediate pol	5.10
	427958	AA418000	Hs.98280	potassium intermediate/small conductance	5.09
	407945	X69208	Hs.606	ATPase, Cu++ transporting, alpha polypep	5.08
20	418576	AW968159	Hs.289104	Alu-binding protein with zinc finger dom	5.05
	413328	Y15723	Hs.75295	guanylate cyclase 1, soluble, alpha 3	5.04
	432729	AK000292	Hs.278732	hypothetical protein FLJ20285	5.Q <u>4</u>
	426342	AF093419	Hs.169378	multiple PDZ domain protein	5.02
~~	429782	NM_005754	Hs.220689	Ras-GTPase-activating protein SH3-domain	5.02
25	436209	AW850417	Hs.254020	ESTs, Moderately similar to unnamed prot	5.02
	430599	NM_004855	Hs.247118	phosphatidylinositol glycan, class B	5.00
	451386	AB029006	Hs.26334	spastic paraplegia 4 (autosomal dominant	5.00
	457211	AW972565	Hs.32399	ESTs, Weakly similar to S51797 vasodilat	4.97
	425851	NM_001490	Hs.159642	glucosaminyl (N-acetyl) transferase 1, c	4.97
30	421689	N87820	Hs.106826	KIAA1696 protein	4.93
	416533	BE244053	Hs.79362	retinoblastoma-like 2 (p130)	4.92
	432653	Ņ62096	Hs.293185	ESTs, Weakly similar to JC7328 amino aci	4.91
	403047	4.F000F00	11- 050500	data (Dunanatana) ilina 4	4.91
25	431117	AF003522	Hs.250500	delta (Drosophila)-like 1	4.90
35	427617	D42063	Hs.199179	RAN binding protein 2	4.88
	428804	AK000713	Hs.193736	hypothetical protein FLJ20706	4.88
	449071	NM_005872	Hs.22960	breast carcinoma amplified sequence 2	4.86 4.84
	407596	R86913	Un 000740	gb:yq30f05.r1 Soares fetal liver spleen	4.84
40	456516	BE172704	Hs.222746	KIAA1610 protein ESTs	4.83
40	458339	AW976853 NM_001141	Hs.172843	arachidonate 15-lipoxygenase, second typ	4.82
	422083 449535	W15267	Hs.111256 Hs.23672	low density lipoprotein receptor-related	4.82
	422048		Hs.288126	spondin 2, extracellular matrix protein	4.82
	424602	NM_012445 AK002055	Hs.151046	hypothetical protein FLJ11193	4.78
45	410765	A1694972	Hs.66180	nucleosome assembly protein 1-like 2	4.77
73	419879	Z17805	Hs.93564	Homer, neuronal immediate early gene, 2	4,74
	450649	NM_001429	Hs.25272	E1A binding protein p300	4.74
	411624	BE145964	Hs.103283	KIAA0594 protein	4.72
	404721	20110001	. 10.110000		4.70
50	426261	AW242243	Hs.168670	peroxisomal famesylated protein	4.70
-	416276	U41060	Hs.79136	LIV-1 protein, estrogen regulated	4.64
	408374	AW025430	Hs.155591	forkhead box F1	4.64
	451900	AB023199	Hs.27207	KIAA0982 protein	4.63
	421437	AW821252	Hs.104336	hypothetical protein	4.63
55	434629	AA789081	Hs.4029	glioma-amplified sequence-41	4.60
	403764				4.58
	421247	BE391727	Hs.102910	general transcription factor IIH, polype	4.53
	403721				4.50
	453070	AK001465	Hs.31575	SEC63, endoplasmic reticulum translocon	4.49
60	417412	X16896	Hs.82112	interleukin 1 receptor, type I	4.48
	439735	AI635386	Hs.142846	hypothetical protein	4.48
•	430261	AA305127	Hs.237225	hypothetical protein HT023	4.46
	430598	AK001764	Hs.247112	hypothetical protein FLJ10902	4.44
	400303	AA242758	Hs.79136	LIV-1 protein, estrogen regulated	4.42
65	438209	AL120659	Hs.6111	aryl-hydrocarbon receptor nuclear transl	4.42
	417421	AL138201	Hs.82120	nuclear receptor subfamily 4, group A, m	4.40
	447270	AC002551	Hs.331	general transcription factor IIIC, polyp	4.38
	434423	NM_006769	Hs.3844	LIM domain only 4	4.35
	404561				4.32

	422969	AA782536	Hs.122647	N-myristoyttransferase 2	4.32
	423685	BE350494	Hs.49753	uveal autoantigen with coiled coil domai	4.32
	425071	NM_013989	Hs.154424	delodinase, iodothyronine, type II	4.32
	431583	AL042613	Hs.262476	S-adenosylmethionine decarboxylase 1	4.31
5	442818	AK001741	Hs.8739	hypothetical protein FLJ10879	4.30
-	423740	Y07701	Hs.293007	aminopeptidase puromydn sensitive	4.24
	424701		Hs.151988	mitogen-activated protein kinase kinase	4.21
		NM_005923			4.20
	424085	NM_002914	Hs.139226	replication factor C (activator 1) 2 (40	4.18
10	410294	AB014515	Hs.323712	KIAA0615 gene product	
10	447124	AW976438	Hs.17428	RBP1-like protein	4.18
	438018	AK001160	Hs.5999	hypothetical protein FLJ10298	4.16
	443857	Al089292	Hs.287621	hypothetical protein FLJ14069	4.15
	446711	AF169692	Hs.12450	protocadherin 9	4.15
	405403				4.14
15	448148	NM_016578	Hs.20509	HBV pX associated protein-8	4.13
	417531	NM_003157	Hs.1087	serine/threonine kinase 2	4.12
	433345	Al681545	Hs.152982	hypothetical protein FLJ13117	4.10
	432712	AB016247	Hs.288031	sterol-C5-desaturase (fungal ERG3, delta	4.09
	435114	AA775483	Hs.288936	mitochondrial ribosomal protein L9	4.08
20	445459	Al478629	Hs.158465	likely ortholog of mouse putative IKK re	4.08
20		M410023	115.130403	tivesh outpoind or monace barranse issue to	4.04
	402791	1100740	Un 0040	Homo sapiens, clone IMAGE:3010666, mRNA,	4.04
	438660	U95740	Hs.6349		
	447568	AF155655	Hs.18885	CGI-116 protein	4.04
0.5	452211	Al985513	Hs.233420	ESTs	4.02
25	443292	AK000213	Hs.9196	hypothetical protein	4.01
	420911	U77413	Hs.100293	O-linked N-acetylglucosamine (GlcNAc) tr	4.00
	428738	NM_000380	Hs.192803	xeroderma pigmentosum, complementation g	3.95
	430456	AA314998	Hs.241503	hypothetical protein	3.95
	437531	Al400752	Hs.112259	T cell receptor gamma locus	3.93
30	428695	Al355647	Hs.189999	purinergic receptor (family A group 5)	3.91
	410011	AB020641	Hs.57856	PFTAIRE protein kinase 1	3.91
	446494	AA463276	Hs.288906	WW Domain-Containing Gene	3.91
	409928	AL137163	Hs.57549	hypothetical protein dJ473B4	3.90
	411598	BE336654	Hs.70937	H3 histone family, member A	3.90
35	425707	AF115402	Hs.11713	E74-like factor 5 (ets domain transcript	3.90
55				RNA 3'-terminal phosphate cyclase	3.89
	451806	NM_003729	Hs.27076	rith 3-termina prospitate cyclase	3.89
	401045	4.4000000	11- 000004	humathatiani amin'n DVE7-547C409	3.89
	437372	AA323968	Hs.283631	hypothetical protein DKFZp547G183	
40	417067	AJ001417	Hs.81086	solute carrier family 22 (extraneuronal	3.88
40	410467	AF102546	Hs.63931	dachshund (Drosophila) homolog	3.88
	431930	AB035301	Hs.272211	cadherin 7, type 2	3.88
	453047	AW023798	Hs.286025	ESTs	3.88
	401785				3.88
	458229	A1929602	Hs.177	phosphatidylinositol glycan, class H	3.86
45	406414				3.86
	412494	AL133900	Hs.792	ADP-ribosylation factor domain protein 1	3.84
	418329	AW247430	Hs.84152	cystathionine-beta-synthase	3.83
	424850	AA151057	Hs.153498	chromosome 18 open reading frame 1	3.82
	427585	D31152	Hs.179729	collagen, type X, alpha 1 (Schmid metaph	3.82
50	423052	M28214	Hs.123072	RAB3B, member RAS oncogene family	3.82
50	416111	AA033813	Hs.79018	chromatin assembly factor 1, subunit A (3.82
	419423	D26488	Hs.90315	KIAA0007 protein	3.80
				FYVE-finger-containing Rab5 effector pro	3.80
	429643	AA455889	Hs.167279		3.80
55	431499	NM_001514	Hs.258561	general transcription factor IIB	
55	444078	BE246919	Hs.10290	U5 snRNP-specific 40 kDa protein (hPrp8-	3.78
	430291	AV660345	Hs.238126	CGI-49 protein	3.76
	431637	A1879330	Hs.265960	hypothetical protein FLJ10563	3.74
	440411	N30256	Hs.151093	hypothetical protein DKFZp434G1415	3.74
	405917				3.74
60	451230	BE546208	Hs.26090	hypothetical protein FLJ20272	3.73
	429597	NM_003816	Hs.2442	a disintegrin and metalloproteinase doma	3.73
	415075	L27479	Hs.77889	Friedreich ataxia region gene X123	3.72
	440351	AF030933	Hs.7179	RAD1 (S. pombe) homolog	3.70
	443603	BE502601	Hs.134289	ESTs. Weakly similar to KIAA1063 protein	3.70
65	446965	BE242873	Hs.16677	WD repeat domain 15	3.70
U.J		Al659306	Hs.73826	protein tyrosine phosphatase, non-recept	3.70
	412350				3.70
	433852	Al378329	Hs.126629	ESTs E1 convers	3.68
	447397	BE247676	Hs.18442	E-1 enzyme	3.68
	405718				3.00

	425217	AU076696	Hs.155174	CDC5 (cell division cycle 5, S. pombe, h	3.68
	421734	Al318624	Hs.107444	Homo sapiens cDNA FLJ20562 fis, clone KA	3.67
	427221	L15409	Hs.174007	von Hippel-Lindau syndrome	3.67
	402408				3.66
5	452946	X95425	Hs.31092	EphA5	3.66
	419078	M93119	Hs.89584	insulinoma-associated 1	3.66
	427144	X95097	Hs.2126	vasoactive intestinal peptide receptor 2	3.65
	423396	Al382555	Hs.127950	bromodomain-containing 1	3.65
	446320	AF126245	Hs.14791	acyl-Coenzyme A dehydrogenase family, me	3.63
10	404939				3.62
	403137				3.60
	437162	AW005505	Hs.5464	thyroid hormone receptor coactivating pr	3.60
	404210				3.59
	443775	AF291664	Hs.204732	matrix metalloproteinase 26	3.56
15	452501	AB037791	Hs.29716	hypothetical protein FLJ10980	3.56
	422443	NM_014707	Hs.116753	histone deacetylase 7B	3.55
	420230	AL034344	Hs.284186	forkhead box C1	3.55
	418428	Y12490	Hs.85092	thyroid hormone receptor interactor 11	3.54
	433002	AF048730	Hs.279906	cyclin T1	3.53
20	405793			•	3.52
	457940	AL360159	Hs.306517	Homo sapiens TRIpartite motif protein ps	3.52
	402444				3.52
	418250	U29926	Hs.83918	adenosine monophosphate deaminase (isofo	3.57
	414222	AL135173	Hs.878	sorbitol dehydrogenase	3.51
25	422384	AA224077	Hs.42438	Sm protein F	3.50
	447805	AW627932	Hs.19614	gemin4	3.50
	454265	H03556	Hs.300949	ESTs, Weakly similar to thyroid hormone	3.50
	423445	NM_014324	Hs.128749	alpha-methylacyl-CoA racemase	3.48
	413435	X51405	Hs.75360	carboxypeptidase E	3.46
30	447210	AF035269	Hs.17752	phosphatidylserine-specific phospholipas	3.46
	426931	NM_003416	Hs.2076	zinc finger protein 7 (KOX 4, done HF.1	3.45
	408418	AW963897	Hs.44743	KIAA1435 protein	3.45
	421887	AW161450	Hs.109201	CGI-86 protein	3.44

Table 7: 42 GENES ENCODING SMALL MOLECULE TARGETS UP-REGULATED IN PROSTATE CANCER COMPARED TO NORMAL ADULT TISSUES

Table 7 shows 42 genes up-regulated in prostate cancer compared to normal adult tissues that are likely to be small molecule targets. These were selected as for Table 5 and the predicted protein contained a structural domain that is indicative of a drugable structure (e.g. protease, kinase, phosphatase, receptor). The functional domain is indicated for each gene.

10	Pkey:

15

Unique Eos probeset identifier number

ExAccn:

Exemplar Accession number, Genbank accession number

UnigenelD:

Unigene number

Unigene Title: PSDomain:

R1:

Unigene gene title Protein Structural Domain Ratio of tumor vs. normal tissue

	Pkey	ExAcen	UnigenelD	Unigene Title	PSDomain	R1	
20	•	AA535210	Un 17100E	kallikrein 3, (prostate specific antigen	trypsin	31.80	
20		X07730		kallikrein 3, (prostate specific antigen	trypsin	24.91	
		X78592	Hs.99915	androgen receptor (dihydrotestosterone r	Androgen_recep,hormone_rec,zf-C4	19.72	
	408430		Hs.44926	dipeptidylpeptidase IV (CD26, adenosine	DPPIV_N_term,Peptidase_S9	16.28	•
		BE245562	Hs.2551	adrenergic, beta-2-, receptor, surface	7tm 1	15.40	
- 25		U80034	Hs.68583	mitochondrial intermediate peptidase	Peptidase_M3	14.81	
		U29589	Hs.7138	cholinergic receptor, muscarinic 3	7tm_1	12.04	
		D50640		phosphodiesterase 3B, cGMP-inhibited	PDEase	11.10	
		U52077	1101007010	gb:Human mariner1 transposase gene, comp		11.02	
	401424	002011		82	arginase	9.58	
30		AB041036	Hs.57771	kallikrein 11	trypsin	9.03	
	428330		Hs.2256	matrix metalloproteinase 7 (matrilysin,	Peptidase_M10	8.76	
	424099	AF071202	Hs.139336	ATP-binding cassette, sub-family C (CFTR	ABC_tran_ABC_membrane	7.64	
	419991	AJ000098	Hs.94210	eyes absent (Drosophila) homolog 1	Hydrolase	7.20	
	431992	NM_002742	Hs.2891	protein kinase C, mu	pkinase,DAG_PE-bind,PH	6.49	
35	447359	NM_012093	Hs.18268	adenylate kinase 5	adenylatekinase	6.00	
	400301	X03635	Hs.1657	estrogen receptor 1	Oest_recep,zf-C4,hormone_rec	5.78	
	421685	AF189723	Hs.106778	ATPase, Ca++ transporting, type 2C, memb	E1-E2_ATPase,Hydrolase	5.37	
	444042	NM_004915		ATP-binding cassette, sub-family G (WHIT	ABC_tran	5.31	
4.0		M73700		lactotransferrin	transferrin,7tm_1	5.29	
40	407945	X69208	Hs.606	ATPase, Cu++ transporting, alpha polypep	E1-E2_ATPase,Hydrolase,HMA	5.08	
	403047				trypsin	4.91	
		D42063		RAN blnding protein 2	Ran_BP1,zf-RanBP,TPR,pro_Isomeras		4.88
		NM_001141		arachidonate 15-lipoxygenase, second typ	lipoxygenase PLAT	4.82	
4.5		W15267	Hs.23672	low density lipoprotein receptor-related	ldl_recept_b,ldl_recept_a,EGF	4.82	
45		NM_013989		deiodinase, lodothyronine, type II	T4_deiodinase	4.32	
	423740			aminopeptidase puromycin sensitive	Peptidase_M1	4.24	
		NM_005923	HS.151988	mitogen-activated protein kinase kinase	pkinase	4.21	
		NM_002914		replication factor C (activator 1) 2 (40	AAA,Viral_helicase1	4.20	
50		NM_003157	Hs.1087	serine/threonine kinase 2	pkinase	4.12	
30		AI355647		purinergic receptor (family A group 5)	7tm_1	3.91	
		AB020641		PFTAIRE protein kinase 1	pkinase .	3.91 3.82	
		AA151057	Hs.73826	chromosome 18 open reading frame 1	Idl_recept_a Y_phosphatase,Band_41,PDZ	3.70	
		Al659306 BE247676	Hs.18442	protein tyrosine phosphatase, non-recept	Hydrolase	3.68	
55	452946		Hs.31092	E-1 enzyme	EPH_lbd.fn3.pkinase,SAM	3.66	
55	452546		Hs.2126	EphA5 vasoactive intestinal peptide receptor 2	7tm_2	3.65	
		AF291664		matrix metalloproteinase 26	Peptidase M10	3.56	
	-	AL360159		Homo sapiens TRIpartite motif protein ps	SPRY,7tm_1	3.52	
	418250		Hs.83918	adenosine monophosphate deaminase (isofo		3.51	
60	413435		Hs.75360	carboxypertidase E	Zn_carbOpept	3.46	
50	-	AF035269	Hs.17752	phosphatidylserine-specific phospholipas	lipase	3.46	
	441210	W 799509	113.17732	huoshuanning-shacing huoshumbas	whase	U.7U	

TABLE 8: 136 GENES SIGNIFICANTLY DOWN-REGULATED IN PROSTATE CANCER COMPARED TO NORMAL PROSTATE

Unique Fos ambeset identifier number

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Table 8 shows 136 genes significantly down-regulated in prostate cancer compared to normal prostate. These were selected from 59680 probesets on the Affymetrix/Eos Hu03 GeneChip array such that the ratio of "average" normal prostate to "average" prostate cancer tissues was greater than or equal to 2. The "average" normal prostate level was set to the mean amongst 4 normal prostate tissues. The "average" prostate cancer level was set to the 85th percentile amongst 73 tumor samples. In order to remove gene-specific background levels of non-specific hybridization, the 10th percentile value amongst all the tissues was subtracted from both the numerator and the denominator before the ratio was evaluated.

	Pkey:			os probeset identifier number					
	ExAccn:								
15	Unigene		Unigene						
	Unigene	e Title:		gene title					
	R1:		Ratio of	normal prostate to prostate cancer					
	Pkey	ExAccn	Cleneninii	Unigene Title		R1			
20	rkey	-	Offigurers	· ·					
20	425932	M81650	Hs.1968	semenogelin I		57.69			
		N98529		Human mRNA for myosin light chain 3 (MLC		19.70			
		X69490	Hs.172004			15.25			
		R41823	Hs.7413	ESTs; calsyntenin-2		10.05			
25		X90568	Hs.172004	titin		9.38			
		D60641	Hs.21739	Homo sapiens mRNA; cDNA DKFZp586i1518 (f		9.05			
		X51501	Hs.99949	protactin-induced protein		8.18			
	411987	AA375975	Hs.183380	*ESTs, Moderately similar to ALU7_HUMAN		7.45			
	404567					5.62			
30	416030	H15261	Hs.21948			5.51			
	444892	Al620617	Hs.148565	ESTs		5 <i>2</i> 7			
	444573	AW043590	Hs.225023	ESTs		5.20			
	428068	AW016437	Hs.233462	ESTs		5.08			
	437440	AA846804	Hs.123694	ESTs		4.95			
35	404113			•		4.75			
			Hs.61260	hypothetical protein FLJ13164		4.75			
			Hs.188181			4.63			
		AV654382	Hs.17947	"ESTs, Weakly similar to K02F3.10 [C.ele		4.53			
40	405163		-	·		4.49			
40	405227					4.45			
		NM_00315		statherin		4.45 4.40			
	450152	Al138635	Hs.22968	ESTs		4.03			
		U35637		"gb:Human nebulin mRNA, partial cds"		4.02			
45	403612		11. 405040	POT-		4.00			
43			Hs.135646			3.98			
				Homo sapiens clone TUA8 Cri-du-chat regi		3.95			
			HS. 128993	"ESTs, Weakly similar to KIAA0465 protei "gb:CM4-HT0244-111199-040-h12 HT0244 Hom		3.95			
		BE148877	11- 470700			3.92			
50				zinc transporter		3.85			
50		AW860972		"gb:QV0-CT0387-180300-167-h07 CT0387 Hom Human clone A9A2BR11 (CAC)n/(GTG)n repea		3.75			
		AF069478	U2'01 120	"gb:AF069478 Homo saplens astrocytoma li		3.61			
	403649	AFU09470		guini cosaro nuttu sapiens astrocytoma ii		3.60			
		H13139	Hs.92282	paired-like homeodomain transcription fa		3.58			
55			Hs.73980	"troponin T1, skeletal, slow"		3.51			
55	414202	AVA130241	Hs.46609			3.45			
	414200	NIM 00020	0Hs.177888	histatin 3		3.37			
	427418	V V 35U 333	Hs.130865	FSTe		3.35			
	42U111 42Q124	AAA21772	Hs.161008	FSTs		3.31			
60		R02018	He 168640	*Ank, mouse, homolog of*		3.30			
00				*EST, Highly similar to ubiquitin-protei		3.30			
	418833	AW074800	Hs.292776	ESTs .		3.26			
		X83957	Hs.83870	nebulin		3.16			
		A00001	, 10.0001 0			-			

	A13778	AA090235	Hs 75535	"myosin, light polypeptide 2, regulatory	3.06
		AW838068	110110000	"gb:QV3-LT0048-010300-109-f02 LT0048 Hom	3.05
			11- 00000		
		AA830811			2.98
	457065	Al476318	Hs.192480	ESTs	2.95
5	432456	H00093		"gb:ph8f12u_19/1TV Outward Alu-primed hn	2.92
-	405678			0.1	2.85
		\$72040	Hs.931	"muncin hagres not montide 2 ekalatal m	2.81
		\$73840		"myosin, heavy polypeptide 2, skeletal m	
		AW189097			2.78
	433968	AL157518	Hs.90421	PRO2463 protein	2.73
10	438522	AAB09431	Hs.258886	ESTs	2.73
		H71937		"complement component 1, s subcomponent"	2.68
		AA102268			2.67
	• •		11342173		2.65
		BE072259		"gb:QV4-8T0536-271299-059-g04 BT0536 Hom	
	415380	F07953	Hs.16085	putative G-protein coupled receptor	2.65
15	428729	AL162331	Hs.191436	hypothetical protein FLJ10619	2.64
		AW207734		"gb:UI-H-BI2-age-h-01-0-UI.s1 NCI_CGAP_S	2.63
			He 152108	transcriptional unit N143	2.63
			113-132-100		2.63
		BE072092		*gb:PM4-BT0532-160200-003-b11 BT0532 Hom	
		M21665	Hs.929	"myosin, heavy polypeptide 7, cardiac mu	2.62
20	437507	AA758538	Hs.246882	ESTS	2.60
	410384	A1933794	Hs.42745	ESTs .	2.58
			Hs.124764		2.58
		AA829828			2.52
					2.51
25				"ESTs, Highly similar to FXD3_HUMAN FORK	
25		Al689154			2.50
	436915	AA737400	Hs.142230	ESTs	2.50
	410028	AW576454	Hs.258553	ESTs	2.46
				alkylglycerone phosphate synthase	2.45
		Al638562		"rp:ts50a10.x1 NCI_CGAP_Ut1 Homo sapiens	2,44
30			Un 400007		2.40
<i>3</i> 0		AA015767			
		H87863	Hs.151380		2.36
	451237	AW600293		*gb:EST00049 pGEM-T library Homo sapiens	2.36
	400001			AFFX control: BioB-3	2.36
		Z45365		"gb:HSC2NF061 normalized infant brain cD	2.36
35		AW872527	Hc 50761		2.36
55					2.36
		AW242394			
		AA742221	Hs.120633		2.35
	407449	AJ002784		gb:Homo sapiens mRNA; fetal brain cDNA 5	2.33
	430573	AA744550	Hs.136345	ESTs	2.32
40	401974				2.31
		AL 044409	He 122262	"ESTs, Weakly similar to PH0217 reverse	2.31
					2.25
				transient receptor potential channel 5	2.25
		Al949371			
	448765	R15337	Hs.21958	"Homo sapiens cDNA FLJ10532 fis, clone N	2.25
45	451130	A1762250	Hs.211347	ESTs	2.24
	405420				2.23
		AW851258		"ab:IL3-CT0220-160200-068-H06 CT0220 Hom	2.23
				"ab:on91f04.s1 Soares NFL_T_GBC_S1 Homo	2.23
		AA933999		· · · · · · · · · · · · · · · · · · ·	
		BE008347		"gb:CM0-BN0154-080400-325-h04 BN0154 Hom	2.23
50		BE252470		gb:601108292F1 NIH_MGC_16 Homo sapiens	2.23
	437010	AA741368	Hs.291434	ESTs	2.23
		Al914279			2.22
	403375				2.21
•		4140000444		Tel-1004 CT0050 020100 022 n00 CT0052 Hom	2.21
EE		AW853441		"gb:RC1-CT0252-030100-023-g09 CT0252 Hom	
55		AW854153		*gb:RC3-CT0254-060400-029-d03 CT0254 Hom	2.20
	421154	AA284333	Hs.287631	"Homo sapiens cDNA FLJ14269 fis, clone P	2.19
	401963				2.18
	435034	AF168711	Hs 159397	x 010 protein	2.18
				KIAA0553 protein	2.18
60					2.17
JU		AW297599			
		Al733395		— · · ·	2.17
	419310	AA236233	Hs.188716	ESTs	2.16
	418579	H91800	Hs.124156	ESTs	2.16
		R54109	Hs.26096		2.16
65					2.15
O)		AA988835			
		AJ133482	rts.165210	COIS	2.15
		AA425562		"gb:zw46e05.r1 Soares_total_fetus_Nb2HF8	2.15
	437101	AA744518	Hs.120610	ESTs	2.15
				"ESTs, Highly similar to collapsin-2-lik	2.15
					-

	415708 459619	H56475		*gb:yt87d11.r1 Soares_pineal_gland_N3HPG	2.13 2.12
		AK000134	Hs 179100	hypothetical protein FLJ20127	2.12
		AA804174	Hs.184354		2.10
5	410881	AW809157		"qb:RC0-ST0118-041099-031-c07_1 ST0118 Homo sapiens cDNA, mRNA sequence"	2.10
•	403087				2.10
	403869			·	2.10
		D81194	Hs.282499	ESTs	2.10
		H29505		"gb:ym60d10,r1 Soares infant brain 1NIB Homo sapiens cDNA clone 5', mRNA sequence"	2.10
10		H11257	Hs.295233		2.09
		BE218221	Hs.190044	ESTs	2.08
	426998	BE274360		"qb:601121068F1 NIH_MGC_20 Homo sapiens cDNA clone 5', mRNA sequence"	2.08
	405455				2.08
	423843	AA332652		"gb:EST36627 Embryo, 8 week I Homo sapiens cDNA 5' end similar to similar to	
15 -				monoamine oxidase B, mRNA sequence"	2.08
	406135				2.07
	427046	BE246180	Hs.121385	ESTs	2.07
	403493				2.05
	444514	A1682905	Hs.270431	*ESTs, Weakly similar to ALU1_HUMAN ALU SUBFAMILY J SEQUENCE	
20				CONTAMINATION WARNING ENTRY [H.sapiens]"	2.05
	435884	AA701443	Hs.192868	ESTs	2.05
	419629	AB020695	Hs.91662	KIAA0888 protein	2.03
	405900			•	2.03
	457350	AW974438	Hs.194136	"ESTs, Moderately similar to AF091457 1 zinc finger protein RIN ZF [R.norvegicus]"	2.02
25	400007			AFFX control: BioDn-5	2.01
	406978	M64358		"gb:Human rhom-3 gene, exon."	2.00

TABLE 8A shows the accession numbers for those primekeys lacking a unigeneID in Table 8. For each probeset we have listed the gene cluster number from which the oligonucleotides were designed. Gene clusters were compiled using sequences derived from Genbank ESTs and mRNAs. These sequences were clustered based on sequence similarity using Clustering and Alignment Tools (DoubleTwist, Oakland California). The Genbank accession numbers for sequences comprising each cluster are listed in the "Accession" column.

10	Pkey: CAT number: Accession:		Unique Eos probeset identifier number Gene cluster number Genbank accession numbers
15	Pkey	CAT number	Accessions
		1014849_1	BE008347 BE008320 BE083307 BE083311 AW075968
		1064753_1	AW207734 D60164 D81150 D81078 D61356 AW996804
20		1154677_1	AW854153 AW500210 BE145772 AW501310
20		1225682_1	AW809157 AW812181 AW812175 AW812172 AW812161 AW812165 AW860972 AW862598 AW862599 AW860988 AW860983 AW860998 AW860925 AW860922 AW860986 AW860984 AW860989
		1256906_1 1353792 1	BE072092 BE072106 BE072086 BE072098 BE072103
		1375933 2	BE252470 BE147573
		1548209_1	H56475 F29401 F34552
25		1558511_1	Z45385 R25905 H06203 T77496
		210744 1	Al638562 T16929 H13401 F07773 R55836
		225415 1	AW838068 AW837986 AW838067 AA322487 AW837936
	423843	232510_1	AA332652 AA331633 AW999369 AW902993 BE170475 AA378845 AW964175 Al475221
	424770	243504_1	AA425562 Al880208 AA346646 N22655 AW811775 AW811786
30	426998	2742591	BE274360
		347718_2	H00093 H00079 H00070 H00054 H00049 H00063 AW905306 AW905241 AW905410 AW905307 AW905411 AW905240
	AW9052	210	A LINE CONTROL OF THE
	400004	450050 4	AW905352 AW905304 AW905239 AW905242 AW905243 H00087
35		452656_1	AA933999 AA761181 H29505 R18575 Z43580 T48738 AI435454 BE004683
33		740749_1 863269_1	AW600293 AI767468
		1249374_1	AW851258 AW851435 AW851106 AW851421
		1251259_1	AW853441 BE145228 BE145218 BE145162 BE145283
		1335127_1	BE072259 BE072230 BE007911
40		543550_1	AF069478 AF069479 AF069480
		_	

TABLE 8B shows the genomic positioning for those primekeys lacking unigene ID's and accession numbers in table 8. For each predicted exon, we have listed the genomic sequence source used for prediction. Nucleotide locations of each predicted exon are also listed.

5							
10	Pkey: Ref: Strand: Nt_position:		Unique number corresponding to an Eos probeset Sequence source. The 7 digit numbers in this column are Genbank Identifier (GI) numbers. "Dunham I. et al." refers to the publication entitled "The DNA sequence of human chromosome 22." Dunham I. et al., Nature (1999) 402:489-495. Indicates DNA strand from which exons were predicted. Indicates nucleotide positions of predicted exons.				
15	Pkey	Ref	Strand	Nt_position			
20	401963	3126783	Plus	51382-51521			
	401974	3126777	Plus	85330-85683			
	403087	8954241	Plus	169511-169795			
	403375	9255944	Minus	92554-92795			
	403493	7341425	Plus	157568-159084			
	403612	8469060	Minus	94723-94859			
	403649	8705159	Minus	27141-27247			
25	403869	7280046	Minus	34379-34583			
	404113	9588571	Minus	13446-13646			
	404567	7249169	Minus	101320-101501			
	405163	9966267	Minus	161171-161299			
	405227	6731245	Minus	22550-22802			
30	405420	7211837	Minus	13428-13582			
	405455	7656675	Plus	134112-134671			
	405678	4079670	Plus	151821-152027			
	405900	6758795	Minus	71181-71535			
	406135	9164918	Minus	65489-65715			

TABLE 9: 1001 GENES SIGNIFICANTLY UP-REGULATED IN NORMAL PROSTATE COMPATED TO PROSTATE CANCER

Table 9 shows 1001 genes significantly up-regulated in prostate cancer compared to normal prostate. These were selected from 59680 probesets on the Affymetrix/Eos Hu03 GeneChip array such that the ratio of "average" normal prostate to "average" prostate cancer tissues was greater than or equal to 8.14. The "average" normal prostate level was set to the mean amongst 4 normal prostate tissues. The "average" prostate cancer level was set to the 85th percentile amongst 73 tumor samples. In order to remove gene-specific background levels of non-specific hybridization, the 10th percentile value amongst all the tissues was subtracted from both the numerator and the denominator before the ratio was evaluated.

5

10

15	Pkey: ExAcon: UnigenelD: Unigene Title: R1:		Unique Eos probeset identifier number Exemplar Accession number, Genbank accession number Unigene number Unigene gene title Ratio of prostate cancer to normal prostate					
20	Pkey	ExAccn	UnigenelD	Unigene Title	R1			
20	451002	AA013299	Hs.8018	ESTs, Weakly similar to ALU3_HUMAN ALU S	1684.00			
		AA689465	Hs.188999		738.00			
		A1078027	Hs.169338		246.86			
		AA928116	Hs.272065		245.20			
25		AK000185		gb:Homo sapiens cDNA FLJ20178 fis, clone	222.00			
	405932				221.33			
		AA864330	Hs.166520	ESTs	212.00			
		AI686550	Hs.174481		163.20			
		Al474866	Hs.193237	ESTs	149.45			
30		NM_002118	Hs.1162	major histocompatibility complex, class	126.11			
		M36860	Hs.9295	elastin (supravalvular aortic stenosis,	123.27			
		AW138330	Hs.233778		120.00			
		X02994	Hs.1217	adenosine deaminase	106.75			
	404407				105.71			
35	442027	A1652926	Hs.128395	ESTs	100.53			
	433704	AA608684		ESTs, Moderately similar to ALUC_HUMAN!	94.00			
	453758	U83527		gb:HSU83527 Human fetal brain (M.Lovett)	89.18			
	415354	F06495		gb:HSC1AB051 normalized infant brain cDN	87.73			
	424239	M67439	Hs.143526	dopamine receptor D5	86.82			
40	444143	AW747996	Hs.160999	ESTs	86.43			
	401672				77.26			
	430590	AW383947	Hs.246381	CD68 antigen	68.47			
	411972	BE074959	•	gb:PM0-BT0582-310100-001-f08 BT0582 Homo	68.00			
	448992	AI766053	Hs.188346		61.26			
45	408828	BE540279		gb:601059857F1 NIH_MGC_10 Homo sapiens c	57.71			
	409653	AW451693	Hs.220826	ESTs	⁻ 56.40			
	402964				54.67			
		N59027		gb:yv59d11.r1 Soares fetal liver spleen	54.00			
~^		AA372275		Homo sapiens cDNA FLJ11383 fis, clone HE	54.00			
50		R32704	Hs.301298	ESTs	52.96			
	405172				52.96			
		AW137088	Hs.144857		52.32			
		AW592931	Hs.256298		51.63			
ب ب		AB028989		mitogen-activated protein kinase 8 inter	50.98			
55		AA703679	Hs.106999	ESTs, Weakly similar to SYT5_HUMAN SYNAP	49.60			
		AA339666		gb:EST44776 Fetal brain I Homo sapiens c	48.90			
		T54095		gb:ya92c05.s1 Stratagene placenta (93722	47.98			
		AA424163	Hs.156895		46.83			
<i>(</i> 0	435672	Al700148	Hs.283626		43.57			
60		AA485224		G protein-coupled receptor kinase-intera	43.00			
		AA837098	Hs.269933		42.70			
	438854	AF074994	Hs.24240	E918	42.67			

	400404				40.42
	406134	4.4.400005	11-004550	FOT- Mississipples to T17000 hypotheti	42.43
		AA480895	HS.201552	ESTs, Weakly similar to T17288 hypotheti	42.31
		AA070266		gb:zm69d04.r1 Stratagene neuroepithelium	42.25
5	401124	41074457	11- 470500	FCT-	41.61 40.00
)		Al371157	Hs.178538		39.64
		AB006628 AW062439	IIS.90400	KIAA0290 protein gb:MR0-CT0060-120899-001-f08 CT0060 Homo	39.60
		AA923278	He 20000E	ESTs, Weakly similar to protease [H.sapl	38.73
		BE221682	Hs.178364		38.06
10		W79114	Hs.58558		36.69
10		AA604799		ESTs, Moderately similar to ALU1_HUMAN A	36.29
		AW963705		ESTs, Weakly similar to ALU7_HUMAN ALU S	36.18
		AA936282	Hs.120397		36.10
		AA333990		coagulation factor XIII, A1 polypeptide	36.08
15		BE314852		hypothetical protein FLJ10257	36.00
	415911	H08796	Hs.124952		36.00
	457502	AA076049	Hs.274415	Homo sapiens cDNA FLJ10229 fis, clone HE	35.23
	421566	NM_000399	Hs.1395	early growth response 2 (Krox-20 (Drosop	35.20
	401468				34.89
20		Al220150	Hs.211195		34.60
		BE350738	Hs.123993	ESTs, Weakly similar to T00366 hypotheti	33.24
		AW848032		gb:lL3-CT0214-231299-053-D11 CT0214 Homo	32.96
	402828				32.93
05		AW518944	Hs.76325	Homo sapiens cDNA: FLJ23125 fis, clone L	31.76
25	402842			CARTION LITORIA III	31.68
		AA285363	H- 4700	gb:HTH280 HTCDL1 Homo sapiens cDNA 5/3	31.59 31.26
		F05183	Hs.1799	CD1D antigen, d polypeptide gb:UI-H-BI1-aea-d-04-0-UI.s1 NCI_CGAP_Su	31.24
		AW139565 H81795		gb:01-7-511-aea-0-04-0-01:51 NCI_CGAP_50 gb:ys68a10.r1 Soares retina N2b4HR Homo	31.20
30		T87479	Hs.291797		31.09
50		AF103907		prostate cancer antigen 3	29.78
		AU076734		solute carrier family 28 (sodium-coupled	29.76
•		Al907039	110.100000	gb:PM-BT134-020499-566 BT134 Homo sapien	29.59
		BE244074	Hs.285531	regulator of Fas-induced apoptosis	29.53
35		AJ870175	Hs.13957		29.47
		R07566	Hs.73817	Small inducible cytokine A3 (homologous	29 .2 2
		W07808		gb:zb03a12.r1 Soares_fetal_lung_NbHL19W	29.20
	448702	AW102670	Hs.122464		29.13
4.0	451939	U80456	Hs.27311	single-minded (Drosophila) homolog 2	28.74
40		W84893	Hs.9305	angiotensin receptor-like 1	28.61
		AB028990		KIAA1067 protein	28.24
		X14008		lysozyme (renal amyloidosis)	28.18
		Al279960	Hs.178140		28.12
45		AW972917		alpha-methylacyl-CoA racemase ESTs, Weakly similar to putative serine/	28.06 27.61
43		AW104257		• •	27.36
	405495	AV650262	NS.75700	GRO2 ancogene	27.33
	406516				27.25
		AW135429	Hs.243577	FSTs	26.96
50		AW452332	Hs.257554		26.36
-		T97490	Hs.50002	small inducible cytokine subfamily A (Cy	26.34
	402838				26.32
		Al979284	Hs.200552		- 26.21
		X57010	Hs.81343	collagen, type II, alpha 1 (primary oste	26.20
55	439792	NM_014856	Hs.6684	KIAA0476 gene product	25.91
	450096	A1682088	Hs.223368		25.60
		AL133660		Homo sapiens mRNA; cDNA DKFZp434M0927 (f	25.57
		BE391090	Hs.280278		25.57
60		NM_005188		Cas-Br-M (murine) ecotropic retroviral t	25.48
60		AA251048	HS.153042	lymphocyte antigen 9	25.42
		AA063426	Un 04400	gb:zf70c08.s1 Soares_pineat_gland_N3HPG	25.25 25.22
		AW083491	Hs.31196	ESTs gb:51f10 Human retina cDNA randomly prim	25.22 25.01
		W28573 T74588	Hs.8509	ESTs, Weakly similar to CO3_HUMAN COMPLE	24.85
65		BE077458	175.000	gb:RC1-BT0606-090500-015-b04 BT0606 Homo	24.76
05		AA760894	Hs.153023	•	24.74
		AI014723	Hs.131770		24.57
		BE019557	Hs.11900	Human DNA sequence from clone RP4-583P15	24.53
		AF026692	Hs.105700	secreted frizzled-related protein 4	24.49
				· · · · · · · · · · · · · · · · · · ·	

					04.40
		U25758	Hs.134584		24.49 24.10
		AL035588		MyoD family inhibitor	24.10
		AA357001	Hs.34045	hypothetical protein FLJ20764	24.04
5		AL122081		cadherin related 23 Homo sapiens cDNA FLJ11720 fis, clone HE	23.89
J		Al208611 AA215672	Hs.12066	gb:zr96e09.s1 NCI_CGAP_GCB1 Homo sapiens	23.83
		AW449674	Hs.47359	ESTs	23.73
		AF204231	Hs.182982		23.62
		AA136301		gb:zk93g04.s1 Soares_pregnant_uterus_NbH	23.39
10		NM_001327	Hs.167379	cancer/testis antigen	23.20
	408380	AF123050	Hs.44532	diublquitin	22.68
	456076	BE243877	Hs.76941	ATPase, Na+/K+ transporting, beta 3 poly	22.65
		AA279530	Hs.83968	integrin, beta 2 (antigen CD18 (p95), ly	22.38
15		R68651	Hs.144997		22.26 22.08
15		BE387335	Hs.283713	KIAA0217 protein	22.04
		AW628686 AW809637	Hs.78851	gb:MR4-ST0124-261099-015-b07 ST0124 Homo	22.00
			Hs.820	homeo box C6	21.95
		AV653846	Hs.126261		21.94
20		BE071874	**********	gb:RC2-BT0522-120200-014-a06 BT0522 Homo	21.84
		J05500	Hs.47431	spectrin, beta, erythrocytic (includes s	21 <i>.</i> 26
	416011	H14487		gb:ym18c10.r1 Soares infant brain 1NIB H	21.24
		Al207936	Hs.7195	gamma-aminobutyric acid (GABA) A recepto	21.14
05		A1623698		Homo sapiens cDNA: FL123529 fis, clone L	21.11
25		X89887	Hs.172350	HIR (histone cell cycle regulation defec	21.10 21.07
		AW502139		gb:Ul-HF-BR0p-ajr-e-05-0-Ul.r1 NIH_MGC_5	20.90
	405685	A1983207	He 192481	ESTs, Weakly similar to SYPH_HUMAN SYNAP	20.84
		AA321355	Hs.285401		20.74
30		AW403724	Hs.140	immunoglobulin heavy constant gamma 3 (G	20.73
	401201				20.73
	458278	W28912	Hs.129019		20.68
		H66948		gb:yr86d10.r1 Soares fetal liver spleen	20.67
25		H42679	Hs.77522	major histocompatibility complex, class	20.66 20.66
35	400926	NIM 004407	Hs.444	serine/threonine kinase 19	20.64
		NM_004197 AW500221	Hs.43616	Homo sapiens mRNA for FLJ00029 protein,	20.61
		X60992	Hs.81226	CD6 antigen	20.61
	405777	7,0000			20.51
40		AW966158	Hs.58582	Homo sapiens cDNA FLJ12702 fis, clone NT	20.20
	425009	X58288		protein tyrosine phosphatase, receptor t	20.10
		BE568568	Hs.195704		19.98
		Al245432		tumor necrosis factor, alpha-induced pro	19.98 19.94
45	418819	AA228776 AA584854	Hs.191721	ab:no09h11.s1 NCI_CGAP_Phe1 Homo sapiens	19.90
43	404426	AA364634		guillosiii 121 NO_OGA _1 net Honio sepiens	19.84
		U43143	Hs.74049	fms-related tyrosine kinase 4	19.79
		NM 012211		integrin, alpha 11	19.62
	414002	NM_006732	Hs.75678	FBJ murine osteosarcoma viral oncogene h	19.57
50	418994	AA296520	Hs.89546	Selectin E (endothelial adhesion molecul	19.56
		AW090198	Hs.4779	KIAA1150 protein	19.52
		AA156781	Hs.83992	ESTs	19.44 - 19.34
		AL138201	Hs.82120	nuclear receptor subfamily 4, group A, m Human pTR7 mRNA for repetitive sequence	19.22
55		X15675 AW449808	Hs.290032	glucosamine (N-acetyl)-6-sulfatase (Sanf	19.21
55		AA284477	Hs.96618		18.77
		Al247422	Hs.129966		18.76
	439845	AL355743	Hs.56663	Homo sapiens EST from clone 41214, full	18.65
	416155	A1807264		ESTs, Weakly similar to AF117610 1 inner	18.64
60	437820	AA769062	Hs.16029		18.62
		AW043951	Hs.38449	ESTs	18.59
		AW247430	Hs.84152	cystathionine-beta-synthase	18.58 18.55
		Al673027 AF113925	Hs.143271		18.52
65	441142 A16061	R42863	Hs.19405 Hs.7124	ESTs	18.47
05	410201 440770	AA912815	Hs.222078		18.40
		A1085846	Hs.25522		18.32
		U51166		thymine-DNA glycosylase	18.28
		AW501751	Hs.279733	ESTs	18.15

		N57568	Hs.176028		18.13
		AF229178		leucine rich repeat and death domain con	18.12
		AW977385	Hs.278615		18.12
_		N90866		CDW52 antigen (CAMPATH-1 antigen)	17.90
5		Al971131		ESTs, Weakly similar to alternatively sp	17.82
		AL050102		DKFZP586F1019 protein	17.82
		Al889114	Hs.195663		17.75 27.75
		AK000596	Hs.3618	hippocalcin-like 1	17.72 17.71
10		AW977724	Hs.75968	thymosin, beta 4, X chromosome	17.67
10	401515	Al097439	Hs.135548	FSTs	17.58
		AL045825	Hs.210197		17.55
		AB001914		paired basic amino acid cleaving system	17.54
	432415		Hs.289014		17.50
15		Al188225	Hs.127462		17.50
		R08003	Hs.188013		17.44
		AA152106	Hs.4859	cyclin L. ania-6a	17.36
	414989	T81668		gb:yd29c04.r1 Soares fetal liver spleen	17.31
	444880	AW118683	Hs.154150		17.30
20	417651	R06874	Hs.268628		17.27
	453457	AL037103		ESTs, Weakly similar to unnamed protein	17.22
		AW452533	Hs.143604		17.22
		M93119		insulinoma-associated 1	17.18
05		BE241624		.CD69 antigen (p60, early T-cell activati	17.14
25	-	AF003522	Hs.250500	delta (Drosophila)-like 1	17.14 17.14
		AW877015	He 150505	gb:QV2-PT0010-250300-096-f12 PT0010 Homo	17.14
		U66468 H08170	Hs.113755	cell growth regulatory with EF-hand doma	17.12
		NM_000361	Hs.2030	thrombomodulin	17.01
30 .		AB032959		KIAA1133 protein	17.00
50 .		AW451157	Hs.181157		16.98
		AA830664	Hs.143974		16.94
		A1940425		gb:CM0-CT0052-150799-024-c04 CT0052 Homo	16.92
	404756				16.91
35	422247	U18244	Hs.113602	solute carrier family 1 (high affinity a	16.90
	420568	F09247	Hs.167399	protocadherin alpha 5	16.88
	443559	AI076765	Hs.269899	ESTs	16.80
	438703	Al803373	Hs.31599		16.78
40		AW845985		gb:RC2-CT0163-200999-002-H08 CT0163 Homo	16.70
40	402895		11- 440404	E 40 ith and distribute of claim as with a trace	16.69 16.68
		NM_006441		5,10-methenyltetrahydrofolate synthetase	16.65
		AW449602 AB002367		ESTs, Moderately similar to NK-TUMOR REC double-cortin and CaM kinase-like 1	16.54
		AW451955	Hs.153065		16.52
45		AW190902		cysteine knot superfamily 1, BMP antagon	16.50
13		R23534	Hs.2730	heterogeneous nuclear ribonucleoprotein	16.50
		AB018319	Hs.5460	KIAA0776 protein	16.40
		AA047854	,	gb:zf49g04.r1 Soares retina N2b4HR Homo	16.32
		A1080042	Hs.180450	ribosomal protein \$24	16.30
50		AA534908	Hs.2860	POU domain, class 5, transcription facto	16.28
	439882	AA847856	Hs.124565	ESTs	16.20
		AW135221	Hs.130812		16.09
		AW796342		gb:PM2-UM0027-230200-002-h02 UM0027 Homo	16.04
		AL049610	Hs.95243		16.04
55		NM_003816	Hs.2442	a disintegrin and metalloproteinase doma	16.02
		Al357412		EST - not in UniGene	16.02 15.94
		BE281591	Hs.222933	hypothetical protein FLJ10511	15.93
		AA055800 AV656098		hypothetical protein FLJ20001	15.86
60		AA076769	113.172002	gb:7B02B10 Chromosome 7 Fetal Brain cDNA	15.85
00	405934	AND UTUS		Sam and to Attentionally to the present about	15.84
		AA622037	Hs.166468	programmed cell death 5	15.84
	416208	AW291168	Hs.41295	ESTs	15.48
	410708	AA534370		Homo sapiens cDNA: FLJ22756 fis, clone K	15.42
65		Al199268		ESTs; Weakly similar to IIII ALU SUBFAMI	15.38
		AW807530		gb:CM0-ST0081-130999-054-d02 ST0081 Homo	15.37
	411507	AW850140		gb:IL3-CT0219-261099-023-D11 CT0219 Homo	15.36
		Al916685	Hs.194601		15.29
	416292	AA179233	Hs.42390	nasopharyngeal carcinoma susceptibility	15.26

	406638	M13861		gb:Human T-cell receptor active beta-cha	15. 2 6
	446686	AW138043	Hs.156307	ESTs	15.25
		Al623511	Hs.118567		15.24
		AW292830			
_		•	Hs.255609		15.22
5		BE147740	Hs.104558		15.22
	409521	BE244854	Hs.159578	Homo sapiens mRNA for FLJ00020 protein,	15.16
	420748	AA279956	Hs.88672	ESTs	15.14
	422583	AA410506	Hs.118578	H.saplens mRNA for ribosomal protein L18	15.14
		AB023185		calcium/calmodulin-dependent protein kin	15.12
10		Al862096	Hs.60640	ESTs	
10			NS.00040		15.12
		BE177778		gb:RC1-HT0598-310300-012-f07 HT0598 Homo	15.12
	445467	Al239832	Hs.15617	ESTs, Weakly similar to ALU4_HUMAN ALU S	15.06
	418305	AW006783	Hs.6686	ESTs	15.03
	402812			•	15.02
15		AA732480	Hs.293581	ESTs	15.00
	400991				15.00
		DESIATOA .	No 70770	Museum mutathus bennamenhamma medala form	
		BE314524	Hs.78776	Human putative transmembrane protein (nm	14.96
		AA460421	Hs.30875	ESTs	14.90
	403683				14.84
20	430315	NM_004293	Hs.239147	guanine deaminase	14.80
	451952	AL120173	Hs.301663		14.72
		J05070		matrix metalloproteinase 9 (gelatinase B	14.69
		BE617135	110.101700	gb:601441677F1 NIH_MGC_65 Homo sapiens c	14.67
			Un 450504		
25		AB021225		matrix metalloproteinase 17 (membrane-in	14.65
25		A1638449	Hs.173031		14.63
	431089	BE041395	Hs.283676	ESTs, Weakly similar to unknown protein	14.60
	459145	A1903354		gb:RC-BT029-100199-117 BT029 Homo sapien	14.55
	449650	AF055575	Hs.297647	ESTs, Moderately similar to calcium chan	14.54
	400952			and the control of th	14.46
30		A1734009	He 197600	ECT ductor (not in UniCons)	14.44
50				EST cluster (not in UniGene)	
		AA905097	Hs.85050	phospholamban	14.42
		Al685464	Hs.292638		14.40
	437210	AA311443	Hs.293563	Homo saplens mRNA; cDNA DKFZp586E2317 (f	14.36
	451900	AB023199	Hs.27207	KIAA0982 protein	14.36
35	445800	AA126419	Hs.301632	ESTs	14.32
		AW945992		immunoglobulin lambda locus	14.31
		AW304028	Hs.300578		14.23
		W57550		Homo sapiens cDNA FLJ13181 fis, done NT	14.22
40		AL049278	Hs.16074		14.22
40		BE242639	Hs.75425	ubiquitin associated protein	14.22
	421913	A1934365	Hs.109439	osteoglycin (osteoinductive factor, mime	14.22
	452712	AW838616		gb:RC5-LT0054-140200-013-D01 LT0054 Homo	14.22
	451468	AW503398	Hs.210047	ESTs	14.16
		Y14443	Hs.88219	zinc finger protein 200	14.14
45		S78187		cell division cycle 25B	14.07
13			Hs.283683		
		AW880709			14.07
		AI815831	Hs.184378		14.05
		AI745649		ESTs, Weakly similar to T00066 hypotheti	14.02
	410020		Hs.728	ribonudease, RNase A family, 2 (liver,	13.98
50	411352	NM_002890	Hs.758	RAS p21 protein activator (GTPase activa	13.98
	429848	AF145439	Hs.225946	chemokine (C-C motif) receptor 9	13.95
		BE159999		gb:QV1-HT0412-270300-123-d10 HT0412 Homo	13.90
	400125	D2.00000		9-1-11 1110112 210000 120 0101110112 110110	- 13.88
		ANNANCOON	No occos	humathatical protein	
55		AW406289	Hs.96593	hypothetical protein	13.85
55		A1479094	Hs.170786		13.80
	422695	AA315158		gb:EST186956 HCC cell line (matastasis t	13.80
	424565	AW102723	Hs.75295	guanylate cyclase 1, soluble, alpha 3	13.78
	458048	H30340	Hs.173705	Homo sapiens cDNA: FLJ22050 fis, clone H	13.78
		AI935400	Hs.217286		13.76
60		AW860158		gb:RC0-CT0379-290100-032-b04 CT0379 Homo	13.75
-	410889		Hs.66744	twist (Drosophila) homolog (acrocephalos	
			F15.00744		13.74
		A1908236		gb:IL-BT166-180399-010 BT166 Homo sapien	13.72
		AW857913		gb:RC0-CT0323-231199-031-b05 CT0323 Homo	13.69
	408364	AW015238	Hs.128453	ESTs	13.67
65	425907	AA365752	Hs.155965		13.62
	402359			•	13.60
	401044				13.53
		AW502498	No 157150	ESTs, Weakly similar to zinc finger prot	13.53
				ESTs	
	423050	AA329648	Hs.23804	LUIS	13.49

	430685	A1690234	Hs 191666	ESTs, Weakly similar to reverse transcri		13.47
				ESTs, Weakly similar to unnamed protein		13.46
		AW578849				
		AW080339	Hs.211911			13.44
	435716	Al573283	Hs.38458			13.44
5	439120	H56389		gb:yt87c03.r1 Soares_pineal_gland_N3HPG		13.43
_	402788			6 -7		13.40
		A A COC AAC	Hs.146278	ECTo		13.40
		AA886446	NS. 140270	E018		
	405411					13.38
	426558	AW188574	Hs.24218	ESTs		13.34
10	453506	AA132818	Hs.110407	ESTs, Weakly similar to coded for by C.		13.33
		AL043004		Human serine/threonine kinase mRNA, part		13.32
				ESTs, Weakly similar to chondroitin 4-su		13.32
		Al074149	ns.130305	ESTS, WEARY SITTING TO CHORDICUM 4-50		
	403838					13.32
	427337	Z46223		Fc fragment of IgG, low affinity IIIb, r		13.30
15	434318	AW207552	Hs.116328	ESTs, Weakly similar to dJ134E15.1 [H.sa		13.28
	435193	N41359	Hs.218107			13.28
		AW451101		ESTs, Moderately similar to hexokinase I		13.27
						13.26
		AF043722		RAS guanyl releasing protein 2 (calclum		
	420052	AA418850	Hs.44410			13.25
20	414020	NM_002984	Hs.75703	small inducible cytokine A4 (homologous		13.25
	403851					13.24
		W07492	Hs.157101	FSTe		13.21
				ESTs, Moderately similar to ALU2_HUMAN A		13.21
		A1762836				
		AB033113	Hs.50187			13.20
25	435063	R21966	Hs.57734	G protein-coupled receptor kinase-intera		13.19
	439367	BE386844	Hs.248746	ESTs		13.17
		A1796320	Hs.10299			13.16
				Homo sapiens cDNA FLJ12334 fis, clone MA		13.14
		AA278362				
~~		BE262802	Hs.4909			13.07
30	426490	NM_001621	Hs.170087	aryl hydrocarbon receptor		13.06
	414789	AA155859	Hs.79708	ESTs		13.05
		BE387790	Hs.26369	ESTs		13.04
		T99719		Homo sapiens cDNA: FLJ22389 fis, clone H		13.03
				ESTs, Weakly similar to putative glycine		13.02
25		AW964806				
35		Al660552		ESTs, Weakly similar to A56154 Abl subst		13.00
	407122	H20276	Hs.31742	ESTs		13.00
	456491	AL137466	Hs.97277	Homo sapiens mRNA; cDNA DKFZp434H1322 (f		12.99
		N75276	Hs.135904			12.98
		AA032197	Hs.102558			12.96
40			Hs.125752			12.96
40		BE267154				
		NM_004354	Hs.79069			12.94
	451154	AA015879	Hs.33536	ESTs		12.93
	412257	AW903830		gb:CM4-NN1037-250400-155-h04 NN1037 Homo		12.93
	449784	AW161319	Hs.12915			12.92
45		D63480		KIAA0146 protein		12.92
TJ						12.92
		NM_001259	Hs.38481			
	439093	AA534163	Hs.5476	serine protease inhibitor, Kazal type, 5		12.90
	416098	H41324	Hs.31581	ESTs, Moderately similar to ST1B_HUMAN S		12.88
	424897	D63216	Hs.153684	frizzled-related protein		12.88
50		AU076649	Hs.76556	growth arrest and DNA-damage-inducible 3		12.88
50		AA587775	Hs.66295	Homo sapiens HSPC311 mRNA, partial cds		12.84
			113.00233	-L. DOC DYCOOD COOLS AND COT DYCEST COO		
		BE077084		gb:RC5-BT0603-220200-013-C07 BT0603 Homo		12.84
	413869	NM_000878	Hs.75596		•	12.80
	452359	BE167229	Hs.29206	Homo sapiens clone 24659 mRNA sequence		12.80
55	435886	BE265839	Hs.12126	hepatocellular carcinoma-associated anti		12.78
		U97018	Hs.12451	echinoderm microtubule-associated protei		12.78
			113.12401	gb:15d7 Human retina cDNA randomly prime		12.77
		W26786		go: 1507 Hullian tenna corex randomy prime		
		AU076643	Hs.313	secreted phosphoprotein 1 (osteopontin,		12.76
	447769	AW873704	Hs.48764	ESTs		12.76
60	414478	Al306389	Hs.76240	adenylate kinase 1		12.76
		D83407		Down syndrome critical region gene 1-lik		12.68
						12.66
		H85157	Hs.40696	ESTs		
	405856					12.66
.	412935	BE267045	Hs.75064	tubulin-specific chaperone c		12.65
65	402802					12.62
		AA889120	Hs.110637	Homeo box A10		12.62
			Hs.93974	and the second s		12.62
		NM_001454	113.333/4	IUINISQU DVA U I		12.60
	403137			to the fact of the contract of		
	430226	BE245562	Hs.2551	adrenergic, beta-2-, receptor, surface		12.57

	448076 AJ1	33123	Hs.20196	adenylate cyclase 9	12.56
	450462 F07	7097	Hs.300828	Homo sapiens mRNA full length insert cDN	12.54
	405236				12.52
	409292 AAI	071051		gb:zm58e05.s1 Stratagene fibroblast (937	12.47
5	421540 AA		Hs.10242	ESTs	12.47
_	425840 AW		Hs.301824	··	12.44
	443181 AIO		Hs.54548		12.42
	452436 BE		Hs.31447	ESTs	12.42
	455183 AW		11001111	gb:RC0-HN0007-160300-011-f09 HN0007 Homo	12.40
10	432887 AI9		Hs.162859		12.37
	410494 M3		Hs.64016	protein S (alpha)	12.36
	439024 R96		Hs.35598	ESTs	12.36
	451246 AW		Hs.39140	cutaneous T-cell lymphoma tumor antigen	12.36
	432892 ALC		Hs.15995	ESTs	
15	418982 AI3		Hs.13073	ESTs	12.35
13	414516 Al3		Hs.279551		12.35
	440134 BE		115.279331	gb:601301619F1 NIH_MGC_21 Homo sapiens c	12.34
	443873 ALC		Hs.16291		12.29
•	401286	M034Z	FIS. 10291	Cols	12.28 12.26
20	454020 AW	negaze	Hs.256527	ECTo	
20	420077 AW		Hs.87767	ESTs	12.24
	420077 AVV		Hs.9884		12.24
			NS.3004	spindle pole body protein	12.24
	407519 X64		Un gener	gb:H.sapiens mRNA HTPCRX01 for oifactory	12.23
25	435839 AF2 448552 AW		Hs.25951	Rho guanine nucleotide exchange factor (12.22
23		9/3033	Hs.20104	hypothetical protein FLJ00052	12.20
	405325	11111	Hs.115707	ECTA	12.20
	451009 AAC				12.18
	423066 Y18		Hs.120171		12.17
30	439556 Al62		Hs.163603		12.16
50	443062 N77		Hs.8963	Homo sapiens mRNA full length insert cDN	12.15
	445873 AA2			Homo sapiens cDNA: FLJ23107 fis, clone L	12.14
	453542 AW		Hs.33190	Homo sapiens mRNA expressed only in plac	12.11
	440106 AA8		Hs.127699		12.10
35	417605 AF0 440286 U29		Hs.82294 Hs.7138	regulator of G-protein signalling 3	12.10
33	420061 AW		Hs.29410	cholinergic receptor, muscarinic 3 ESTs	12.04
	458727 AIO		Hs.92679		12.02
	445407 AI22			Homo sapiens clone CDABP0014 mRNA sequen ESTs, Weakly similar to la costa [D.mela	11.96
	418250 U29		Hs.83918	adenosine monophosphate deaminase (isofo	11.95
40	414129 Al99		Hs.270798		11.94
70	409799 D11		Hs.76845		11.93
				phosphoserine phosphatase-like	11.92
	438461 AW0 443912 R37		Hs.184780	phosphoserine aminotransferase	11.92
	424606 AA3		115.104700	gb:EST49786 Gall bladder I Homo sapiens	11.92
45	434217 AW		Hs.23349	ESTs	11.90
13	451533 NM		Hs.26530	serum deprivation response (phosphatidy)	11.90
	422423 AF2			CD72 antigen	11.90 11.89
	409398 AW		115.110401	gb:PM4-PT0019-121299-004-F02 PT0019 Homo	11.89
	423853 AB0		He 133/66	slit (Drosophila) homolog 1	11.82
50	446180 Al07		Hs.14220	hypothetical protein FLJ20450	11.80
-	414341 D80			KIAA0182 protein	11.80
	406538	001	110.70000	Niroto toz protesti	11.79
	433253 AW	150502	Hs.24218	FSTe	- 11.79
	447397 BE2			E-1 enzyme	11.78
55	451684 AF2			CDA14	11.76
55	416862 R23		Hs.23575		11.74
	425770 NM_			spastic ataxia of Charlevoix-Saguenay (s	11.72
	428826 ALO	18842	Hs.194019	attraction	11.72
	433037 NM_			HSPC067 protein	11.72
60	447476 BE2			ESTs	11.72
50	452092 BE2			hypothetical protein FLJ11210	11.72
	432032 BE2			H2.0 (Drosophila)-like homeo box 1	11.72
	412922 MOU 401680 NM			LIM domain-containing preferred transloc	
	401680 NM_ 422576 BE5	10EEE	He 110030	CGL93 protois	11.69
65	422576 BE5 450203 AF0			CGI-83 protein L-kynurenine/alpha-aminoadipate aminotra	11.68
5 5	450203 AF0		10201020	gb:QV0-CT0224-261099-035-g02 CT0224 Homo	11.68
	410531 AVV7 425917 W28			Homo sapiens cDNA: FLJ23067 fis, clone L	11.67
	423917 W26 418693 Al75			thrombospondin 1	11.66
	410093 AI75	0010	10.01403	anomoosponum i	11.64
	400007				11.62

	416188	BE157260	Hs.79070	v-myc avian myelocytomatosis viral oncog	11.60
		AW952771	Hs.90043	ESTs	11.59
		AI986160	Hs.88446	ESTs	11.59
,,,	400885				11.57
5		AW502327		gb:UI-HF-BROp-aka-a-07-0-UI.rl NIH_MGC_5	11.56
	400802	NII 4 040045	11	Till describle bounds	11.56
		NM_016045	Hs.5184	TH1 drosophila homolog	11.55
		M55994		tumor necrosis factor receptor superfami	11.55 11.54
10		S55736 AA460479	Hs.4096	ESTs, Weakly similar to hypothetical pro KIAA0742 protein	11.53
10		Z42047		ESTs; KIAA0738 gene product	11.52
		AW964897	Hs.290825		11.52
		AA426080	Hs.98489		11.50
		AW204232	Hs.279522		11.50
15		X72755	Hs.77367		11.46
		F18572	Hs.22978	, ,	11.44
		AA453208	Hs.28726		11.42
	459029	AA131376	Hs.285203	fibroblast growth factor 12	11.42
	456267	Al127958		cystatin E/M	11.39
20	433285	AW975944	Hs.237396	ESTs	11.38
	449186	AW291876	Hs.196986		11.37
		A1434593	Hs.164294		11.37
		R00028		gb:ye70a06.s1 Soares fetal liver spleen	11.36
05		A1277652	Hs.54578	ESTs	11.31
25	401163		11-040440	about the 10 mate VO accepted	11.31
		£36149	HS.248116 Hs.47289	chemokine (C motif) XC receptor 1 ESTs	11.28 11.28
		AW246803 AL044829	Hs.29331		11.27
		NM_014253	Hs.23796		11.26
30		AA075687		epidermal growth factor receptor substra	11.24
20		W07411		ESTs, Moderately similar to ALU3_HUMAN A	11.24
		H28383		gb:yl52c03.r1 Soares breast 3NbHBst Homo	11.24
		AA631047	Hs.158761	Homo sapiens cDNA FLJ13054 fis, clone NT	11.23
	442501	AA315267	Hs.23128	ESTs	11.22
35	429746	AJ237672	Hs.214142	5,10-methylenetetrahydrofolate reductase	11.21
	422858	R35398		gb:yg64g10.r1 Soares infant brain 1NIB H	11.20
		X84908	Hs.78060		11.20
		AV660122	Hs.282675		11.20
40		C21322	Hs.11577		11.20
40		W78902	Hs.293297		11.17 11.16
		AI367347	Hs.127809 Hs.214410		11.16
		AW748078 BE142098	N3.2 144 IU	gb:CM4-HT0137-220999-017-d11 HT0137 Homo	11.14
		AB020725	Hs 58009	KIAA0918 protein	11.14
45	405601	ADOLUIZO	1 10.0000	the secto protein	11.13
		Al000341	Hs.220491	ESTs	11.12
		AA410183	Hs.137475		11.12
		N77624	Hs.173717	phosphatidic acid phosphatase type 2B	11.10
·	431475	AI567669	Hs.287316	ESTs	11.10
50		AF030880		solute carrier family, member 4	11.08
		AW104057	Hs.19193	ESTs	11.07
		Y00093	Hs.51077	integrin, alpha X (antigen CD11C (p150),	11.07
		W92745	Hs.193324		- 11.03
55		U52077	11- 40404	gb:Human mariner1 transposase gene, comp	11.02
23		AF055581	Hs.13131	tymphocyte adaptor protein gb:MR1-SN0033-120400-002-c10 SN0033 Homo	11.02 10.95
		AW867079 BE382701	Hs.25960	v-myc avian myelocytomatosis viral relat	10.95
		AW006969	Hs.6311	hypothetical protein FLJ20859	10.94
		AW591783	Hs.36131	collagen, type XIV, alpha 1 (undulin)	10.94
60		AA530994		ghrelin precursor	10.92
		AW246428	Hs.75355	ubiquitin-conjugating enzyme E2N (homolo	10.92
	400132			, , , , , , ,	10.92
•		AA443966	Hs.31595	ESTs	10.90
		NM_000328		retinitis pigmentosa GTPase regulator	10.88
65		D85782	Hs.3229	cysteine dioxygenase, type I	10.88
		Al366213	Hs.173422	KIAA1605 protein	10.87
		AW948126		gb:RC0-MT0013-280300-031-a12 MT0013 Homo	10.85
	400615		14. 481400	FIF to annual time to store t	10.80
	425018	BE245277	Hs.154196	E4F transcription factor 1	10.80

					40 70
	456011	BE243628		gb:TCBAP1D1053 Pediatric pre-B cell acut	10.79
	455982	BE176862		gb:RC4-HT0587-170300-012-a04 HT0587 Homo	10.74
	450418	BE218418	Hs.201802		· 10.73
		AW803564	Hs.288850		10.72
5			Hs.5364	DKFZP564I052 protein	10.70
)		AW377314			
		Al383497		ESTs, Weakly similar to ALU1_HUMAN ALU S	10.70
		R40978		ESTs, Moderately similar to ALU1_HUMAN A	10.70
	449590	AA694070	Hs.268835	ESTs	10.68
	446035	NM_006558	Hs.13565	Sam68-like phosphotyrosine protein, T-ST	10.68
10		U24578	Hs.170250	complement component 4A	10.66
10		AW863261		ESTs, Highly similar to AF161358 1 HSPC0	10.64
					10.64
		AA220238	Hs.94986		
		AF151879	Hs.26706	•	10.62
	438893	AF075031	Hs.29327	ESTs	10.62
15	459324	AW080953		gb:xc28c12.x1 NCI_CGAP_Co18 Homo sapiens	10.61
	439883	AL359652	Hs.171096	Homo sapiens EST from clone DKFZp434A041	10.58
		AA715328	Hs.291205		10.57
		AA128423	Hs.40300		10.57
			Hs.90998		10.56
20		D50918			
20		R10184		ESTs, Weakly similar to ALU1_HUMAN ALU S	10.56
		Al142350	Hs.146735		10.55
	411448	AA178955	Hs.271439	ESTs	10.54
	410102	AW248508	Hs.279727	ESTs;	10.52
	406577			·	10.52
25		AK001332	Hs.44672	hypothetical protein FLJ10470	10.51
		AF059214		cholesterol 25-hydroxylase	10.50
		AT 0032 14	113.134007	Gloid Steror 20-try Gloxy Goo	10.48
	400880	4 4 0 0 1 0 10	11- 5007		
		AA894876	Hs.5687	protein phosphatase 1B (formerly 2C), ma	10.48
	434715	BE005346	Hs.116410		10.46
30	406851	AA609784	Hs.180255	major histocompatibility complex, class	10.44
	413409	AI638418	Hs.21745	ESTs	10.44
		U76421	Hs.85302	adenosine deaminase, RNA-specific, B1 (h	10.44
		AW500239	Hs.21187	Homo sapiens cDNA: FLJ23068 fis, clone L	10.44
		Al909154	110.21107	gb:QV-BT200-010499-007 BT200 Homo sapien	10.44
35		Y18418	He 272822	RuvB (E coli homolog)-like 1	10.44
23				ESTs, Weakly similar to ALU1_HUMAN ALU S	10.42
		R08950			
		AA788946	Hs.16869		10.41
		NM_003878	Hs.78619		10.41
40		NM_003500	Hs.9795	acyl-Coenzyme A oxidase 2, branched chai	10.40
40 ·	458873	AW150717		STAT induced STAT inhibitor 3	10.38
	415082	AA160000	Hs.137396	ESTs	10.37
	429124	AW505086	Hs.196914	minor histocompatibility antigen HA-1	10.36
	417187	AB011151	Hs.81505	KIAA0579 protein	10.34
		AW067805		methylenetetrahydrofolate dehydrogenase	10.34
45		NM_000030	He 271366	alanine-glyoxylate aminotransferase homo	10.33
43		_		ESTs	10.32
		T93096	Hs.17126		
		NM_014324		alpha-methylacyl-CoA racemase	10.31
		AW860597	Hs.30164	ESTs	10.30
	432242	AW022715	Hs.162160	ESTs, Weakly similar to ALU4_HUMAN ALU S	10.30
50	406394	AA172106	Hs.110950	Rag C protein	10.30
	406189				10.29
		AW411307	Hs.114311	CDC45 (cell division cycle 45, S.cerevis	10.26
		AA172106		Rag C protein	- 10.26
		T89832	Hs.170278		10.26
55		NM_006762		Lysosomal-associated multispanning membr	10.24
JJ					
		NM_005211	HS.174142	colony stimulating factor 1 receptor, fo	10.24
	401384				10.23
		D13168	Hs.82002	endothelin receptor type B	10.22
	426928	AF037062	Hs.172914	retinol dehydrogenase 5 (11-cisand 9-cis	10.21
60	423032	A1684746	Hs.119274		10.20
	436556	Al364997		ESTs	10.20
		BE243026		KIAA0246 protein	10.19
	437401	AA757196	Hs.121190		10.19
	403690		. 15. 12 1 100	2010	10.13
65		BE152393		gb:CM2-HT0323-171199-033-a08 HT0323 Homo	10.17
UJ			No googe		
		AA305599		hypothetical protein PRO2013	10.16
		AW975009	Hs.292274		10.16
	432827	Z68128	Hs.3109	Rho GTPase activating protein 4	10.16
	432660	Al288430	Hs.64004	ESTs	10.14

	452234	AW084176	Hs.223296	ESTs		10.14
		Al245701		gb:qk31f05.x1 NCI_CGAP_Kid3 Homo sapiens		10.13
		AA626142		ESTs, Weakly similar to KPCE_HUMAN PROTE		10.13
5		A1174603		enolase 1, (alpha)		10.12
5		A1038989 NM_006056	NS.24009	hypothetical protein FLJ10826		10.12 10.12
		AA807346	Hs 288581	Homo sapiens cDNA FLJ14296 fis, clone PL		10.12
		AW118822	Hs.128757			10.10
		AW014605	Hs.179872			10.10
10	417308	H60720	Hs.81892	KIAA0101 gene product		10.09
		Al204266	Hs.179303			10.05
		A1433833		ESTs, Weakly similar to ALU1_HUMAN ALU S		10.04
		BE614599		H.sapiens gene from PAC 42616, similar t		10.04
15		BE552368 AA329796	Hs.1098	Homo sapiens cDNA FLJ13445 fis, clone PL DKFZp434J1813 protein		10.04 10.02
13		AW206373	Hs.16443			10.00
		X58528	Hs.76781			10.00
	421832	NM_016098	Hs.108725	HSPC040 protein		10.00
••	423554	M90516	Hs.1674	glutamine-fructose-6-phosphate transamin		10.00
20		Al922988	Hs.172510			10.00
		AW137442	Hs.136965			10.00
		AA418280 BE501815	Hs.180040	Homo saplens cDNA: FLJ22439 fis, done H		10.00 9.99
		AA425310	Hs.155766			9.98
25		Al147652		Homo sapiens clone HH409 unknown mRNA		9.98
		N72394	Hs.44862			9.96
	427509	M62505	Hs.2161	complement component 5 receptor 1 (C5a I		9.96
		AB028945		cortactin SH3 domain-binding protein		9.96
20		AW009605	Hs.231923			9.96
30		AW474513 AA704703	Hs.224397 Hs.77031	ESTs, Weakly similar to B48013 proline-r Sp2 transcription factor		9.94 9.94
	434596		113.77031	gb:yb65g12.s1 Stratagene ovary (937217)		9.94
		BE276115	Hs.144980	ESTs, Weakly similar to CA13_HUMAN COLLA		9.93
		AA033813		chromatin assembly factor 1, subunit A (9.92
35	423349	AF010258		homeo box A9		9.92
		AW975531		minichromosome maintenance deficient (S.		9.92
		AW192307		dolichyl-P-Gic:Man9GicNAc2-PP-dolichylgl		9.90
		AA481003 D87450	Hs.97128	KIAA0261 protein		9.90 9.90
40		AW976507	Hs.293515			9.90
		AW972187		hypothetical protein FLJ22215		9.89
	-	NM_005291		G protein-coupled receptor 17		9.88
		Al097570	Hs.71222			9.87
15		AW801383		H.sapiens mRNA for ribosomal protein L18		9.86
45		Al278802	Hs.25661 Hs.245484	ESTS		9.85 9.85
		AW117416 AL043002		ESTs, Moderately similar to unnamed prot		9.84
		Al962552	Hs.226765			9.84
		Al267652	Hs.30504	Homo sapiens mRNA; cDNA DKFZp434E082 (fr		9.82
50	431066	AF026273	Hs.249175	Interleukin-1 receptor-associated kinase		9.82
		AW894667		chimerin (chimaerin) 1		9.80
		A1792888	Hs.145489			9.80
		AL050072 D13635	He 155297	gb:Homo sapiens mRNA; cDNA DKFZp566E1346 KIAA0010 gene product	•	9.75 9.74
55		N67619	Hs.43687			9.74
55		AI590401	Hs.21213			9.73
		Al381659	Hs.267086			9.72
	437672	AW748265	Hs.5741	flavohemoprotein b5+b5R		9.72
CO		AL120659	Hs.6111	KIAA0307 gene product		9.72
60		AA807228	Hs.225161			9.72
		AA311152		ESTs; Weakly similar to KIAA0226 [H.sapi ESTs		9.72
		Al289619 AK001451	Hs.13040	CD2-associated protein		9.72 9.70
	436296			Homo saplens cDNA: FLJ21084 fis, clone C		9.70
65		AW977540	Hs.269254			9.70
		S67580	Hs.1645	cytochrome P450, subfamily IVA, polypept		9.67
		AW118645	Hs.160004			9.67
		Al393657	Hs.159750			9.66
	421044	AF061871	HS.101302	collagen, type XII, alpha 1		9.66

		BE466863	Hs.280099		9.66
		R91679	Hs.124981		9.66
	433480			immunoglobulin lambda locus	9.65
_		Al248301	Hs.127112	= : =	9.65
5		D53304	Hs.65394		9.65
		R27496	Hs.1378	annexin A3	9.64
		C05569		hypothetical protein FLJ13057 similar to	9.64
		AA488101 AA007534	Hs.125062	inactivation escape 1	9.62 9.62
10		AW063190	Hs.279101		9.61
10	406554	M11003130	115.275101	2019	9.60
		AA377607	Hs.273138	FSTs	9.58
		AW392394		KIAA0064 gene product	9.58
		AK001578	Hs.24129		9.58
15		Al494332	Hs.196963		9.58
		AL117474	Hs.41181		9.56
	446572	AV659151	Hs.282961		9.56
	459245	BE242623	Hs.31939	manic fringe (Drosophila) homolog	9.55
	423545	AP000692	Hs.129781	chromosome 21 open reading frame 5	9.54
20	414697	BE266134	Hs.76927		9.54
		AW807057		gb:MR4-ST0062-031199-018-b03 ST0062 Hamo	9.52
		NM_005574		LIM domain only 2 (rhombotin-like 1)	9.52
		D26067		KIAA0033 protein	9.52
25		NM_004573	Hs.994	phospholipase C, beta 2	9.51
23		AW295389	Hs.119768	Homo sapiens cDNA: FLJ22199 fis, clone H	9.51 9.50
		AA742181 D28459		ubiquitin-conjugating enzyme E2A (RAD6 h	9.50
		AA094538	Hs.6588	ESTs	9.50
		AA833902	Hs.270745		9.48
30		R07114	Hs.271224		9.48
		AJ132085		gb:Homo sapiens mRNA for axonemal dynein	9.44
•	458723	AW137726	Hs.244352	ESTs, Moderately similar to laminin alph	9.44
	457526	AW450584	Hs.192131	ESTs, Weakly similar to RIBB [H.sapiens]	9.43
	404741		•		9.43
35		NM_005428	Hs.116237	vav 1 oncogene	9.43
	403708	*****		III I DANA EL IOCEDO Co I O	9.42
		AW847814	HS.289005	Homo sapiens cDNA: FLJ21532 fis, clone C	9.42
	417380		Hs.144967	gb:EST04698 Fetal brain, Stratagene (cat	9.42 9.42
40		AA354690 AA004410		acyl-Coenzyme A oxidase 1, palmitoyl	9.42
40		AU076606		coagulation factor V (proaccelerin, labi	9.42
		AW893569	110.00001	gb:RC0-NN0021-040400-021-c10 NN0021 Homo	9.41
		AA361623	Hs.288775	Homo sapiens cDNA FLJ13900 fis, clone TH	9.41
		AW968504		CDC2-related protein kinase 7	9.40
45	414171	AA360328	Hs.865	RAP1A, member of RAS oncogene family	9.40
	415947		Hs.78934		9.40
		BE262745		gb:601153869F1 NIH_MGC_19 Homo sapiens c	9.39
		A1689987		ESTs, Weakly similar to RMS1_HUMAN REGUL	9.39
50		BE514362	MS.296422	FK506-binding protein 3 (25kD)	9.39
J U	402835				9.38 9.38
	404632 446566	H057/1	Hs.17914	Homo sapiens cDNA: FLJ22801 fis, clone K	9.37
		AW903533	113.17317	ab:CM1-NN1031-060400-178-d05 NN1031 Homo	- 9.37
		Al095087	Hs 152299	ESTs, Moderately similar to ALU5_HUMAN A	9.36
55		Al420611	Hs.127832		9.36
		BE258532		CTP synthase	9,34
	429327	AA283981	Hs.199248	prostaglandin E receptor 4 (subtype EP4)	9.33
	407061	X97748		gb:H.sapiens PTX3 gene promotor region.	9.33
<i>-</i> 0		BE616731	Hs.80645	interferon regulatory factor 1	9.33
60		AW875443	Hs.22209	secreted modular calcium-binding protein	9.33
		AA693960	Hs.103158	·	9.33
		BE513731	Hs.88959	Human DNA sequence from clone 967N21 on	9.32
		AA033699	Hs.83938	ESTs, Moderately, similar to MASP-2 [H.sa	9.32
65		NM_007274	Hs.8679	cytosolic acyl coenzyme A thioester hydr Homo sapiens cDNA: FLJ23591 fis, clone L	9.32 9.32
UJ	452859	Al300555	113.200130	Tionio adpiana contra i cacada i ila, Golia L	9.32
		AW025529	He 230812	ESTs, Weakly similar to CALM_HUMAN CALMO	9.31
		AW976410		Homo sapiens cDNA: FLJ21016 fis, clone C	9.30
	419066		Hs.6975	PRO1073 protein	9.30

		AW167128	Hs.231934	ESTs	9.30
	405125	AMMADOFCO		abd II HE DOG all b co o III at AULI MOO E	9.30
		AW499566	Un E4000	gb:UI-HF-BR0p-aji-h-03-0-UI.r1 NIH_MGC_5	9.28
5		AI191811 AF000652	Hs.54629 Hs.8180	ESTs syndecan binding protein (syntenin)	9.28 9.27
3		AJ250839	Hs.58241		9.26
		AW013907		ESTs, Moderately similar to predicted us	9.26
		AF164142	Hs.82042		9.25
		D29642	Hs.1528	KIAA0053 gene product	9.25
10		AA281279	Hs.23317		9.24
	435980	AF274571		ESTs; Weakly similar to DEOXYRIBONUCLEAS	9.24
	458530	BE395035	Hs.199889	ESTs, Weakly similar to KIAA0874 protein	9.24
	402585				9.24
15		AA280700	11- 400004	gb:zs95h11.s1 NCL_CGAP_GCB1 Homo sapiens	9.23
15		AA431791	Hs.183001		9.22 9.22
		U42349 AW582962		Putative prostate cancer tumor suppresso ESTs, Highly similar to AF151805 1 CGI-4	9.20
		BE514514		coronin, actin-binding protein, 1A	9.19
		AL039185		thyroid hormone receptor interactor 7	9.18
20		Al206589	Hs.116243		9.17
		U61412	Hs.51133		9.17
	428242	H55709	Hs.2250	leukemia inhibitory factor (cholinergic	9.16
		AW080356		ESTs, Weakly similar to alternatively sp	9.15
25		BE182082	Hs.246973		9.14
25		AA528402	Hs.74861		9.14
		AW151660 U54727	Hs.31444 Hs.191445		9.14 9.14
		Al911527	Hs.11805		9.14
		BE075281	110.11000	gb:PM1-BT0585-290200-005-d07 BT0585 Homo	9.12
30	413450		Hs.75372		9.12
-	446442	BE221533	Hs.257858		9.12
	438540	AA810021	Hs.136906	ESTs	9.12
		M24283		Intercellular adhesion molecule 1 (CD54)	9.11
25		AA402307		ubiquinol-cytochrome c reductase hinge p	9.10
35		AA913736	Hs.126715		9.10
		NM_014720 AF086534		Ste20-related serine/threonine kinase ESTs, Moderately similar to ALU1_HUMAN A	9.10 9.10
		C19035	Hs.164259		9.09
		M88700		dopa decarboxylase (aromatic L-amino aci	9.08
40		AW190920	Hs.19928		9.08
	425844	T68073	Hs.159628	serine (or cysteine) proteinase inhibito	9.08
		Al624436	Hs.194488		9.07
		BE328153	Hs.240087		9.06
45	436490		Hs.18713		9.05
47		AA065131 BE300091	He 110600	ESTs, Weakly similar to ALU7_HUMAN ALU S hypothetical protein FLJ12969	9.05 9.04
		AW369351		Homo sapiens cDNA FLJ13090 fis, done NT	9.04
		Al174947		Homo sapiens mRNA; cDNA DKFZp564D1164 (f	9.04
		AW301344	Hs.195969		9.04
50		AU076730		kinesin 2 (60-70kD)	9.02
	418650	BE386750	Hs.86978	prolyl endopeptidase	9.02
		M81590	Hs.123016	5-hydroxytryptamine (serotonin) receptor	9.02
		AA326108	Hs.53631	ESTs	- 9.02
55		BE622585	Hs.3731	ESTs adenosine A2b receptor pseudogene	9.02 9.01
JJ	443453	AW572659	Hs.269882		9.01
		AW972330		triggering receptor expressed on myeloid	9.01
		AW295840		Homo sapiens cDNA: FLJ21513 fis, clone C	9.00
		AB008681	Hs.23994	activin A receptor, type IIB	9.00
60		AA902386	Hs.286	ribosomal protein L4	8.99
		AW341130		ESTs, Moderately similar to FGFE_HUMAN F	8.99
		AF051428		estrogen receptor 2 (ER beta)	8.97
		AA883316	Hs.255221		8.96
65		BE000707	Hs.29567		8.96
UJ		AI051313 BE614387	Hs.143315 Hs.47378		8.96 8.96
		Al084125		transcription factor	8.95
	438707		Hs.5326	porcupine	8.95
	402240			# T 1114 T	8.95

	444152	Al125694	Hs.149305	Homo sapiens cDNA FLJ14264 fis, done PL	8.95
	409842	AW501756	•	gb:UI-HF-BR0p-ajm-c-09-0-UI.r1 NIH_MGC_5	8.94
	416277	W78765	Hs.73580	ESTs	8.94
	456697	Al908006	Hs.111334	ferritin, light polypeptide	8.94
5	410762	AF226053	Hs.66170	HSKM-B protein	8.92
_		AL120344	Hs.75074		8.92
		Al287817	Hs.129636	•	8.92
		AA002064	Hs.18920		8.91
	411486			eukaryotic translation elongation factor	8.90
10		BE566249	Hs.20999	Homo sapiens cDNA: FLJ23142 fis, clone L	8.90
10		AA257161	Hs.8658	hypothetical protein DKFZp434E0321	8.89
		NM 007019	Hs.93002	ubiquitin carrier protein E2-C	8.89
			113.00002	gb:lL3-CT0215-020300-090-E06 CT0215 Homo	8.89 .
		AW849292 AW971228	N° 3003E0		8.89
15		D00723	Hs.290259 Hs.77631	glycine cleavage system protein H (amino	8.88
13			Hs.83337		8.87
		Z37976	IIS.0333/	latent transforming growth factor beta b	8.87
	406422	NII 040400	11- 404740	des financiamists 10	
		NM_016102		ring finger protein 16	8.87
20		D50030	Hs.104	HGF activator	8.86
20	418203			CDC28 protein kinase 2	8.86
		AA744529		mitogen-activated protein kinase kinase	8.85
		H66566	Hs.271711	ESTs	8.85
		AA076049	Hs.274415	Homo sapiens cDNA FLJ10229 fis, clone HE	8.84
		Al952797		.Homo sapiens cDNA: FLJ21559 fis, clone C	8.83
25		T89839	Hs.119471		8.83
	425694	U51333		hexokinase 3 (white cell)	8.82
	419972	AL041465	Hs.294038	ESTs, Moderately similar to ALU2_HUMAN A	8.82
		Al683487		Homo saptens cDNA FLJ11441 fis, done HE	8.82
	413413	D82520	Hs.301834	Homo sapiens cDNA FLJ10952 fis, clone PL	8.82
30		AA435997	Hs.104930	ESTs	8.82
	415839	R40611	Hs.137565		8.81
		N34145	Hs.250614	ESTs	8.80
	420309	AW043637	Hs.21766	ESTs	8.80
	421863	Al952677	Hs.108972	Homo sapiens mRNA; cDNA DKFZp434P228 (fr	8.80
35	447965	AW292577	Hs.94445		8.80
•	459172	BE063380	•	gb:PM0-BT0275-291099-002-g10 BT0275 Homo	8.80
	403259				8.78
	411534	AW850473		gb:IL3-CT0219-280100-061-B11 CT0219 Homo	8.78
	456161	BE264645	Hs.282093	Homo sapiens cDNA: FLJ21918 fis, clone H	8.77
40	413654	AA331881	Hs.75454	peroxiredoxin 3	8.76
	401744				8.76
	425348	AL137477		cadherin-like 24	8.76
	423396	AI382555	Hs.127950	bromodomain-containing 1	8.75
	450649	NM_001429		Human DNA sequence from clone RP1-85F18	8.75
45	408331	NM_007240	Hs.44229	dual specificity phosphatase 12	8.74
	423872	AB020316	Hs.134015	uronyl 2-sulfotransferase	8.74
	424906	A1566086	Hs.153716	Homo sapiens mRNA for Hmob33 protein, 3'	8.74
	427596	AA449506	Hs.179765	Homo sapiens mRNA; cDNA DKFZp586H1921 (f	8.73
	432488	AA551010	Hs.216640		8.72
50	448980	AL137527	Hs.22703	Homo sapiens mRNA; cDNA DKFZp434P1018 (f	8.72
	429455	Al472111	Hs.292507	ESTs	8.71
	429855	AW385597	Hs.138902	ESTs, Weakly similar to B34087 hypotheti	8.71
	441746	H59955	Hs.127829		- 8.70
	411945	AL033527	Hs.92137	v-myc avian myelocytomatosis viral oncog	8.70
55	413492	D87470	Hs.75400	KIAA0280 protein	8.70
		W31254	Hs.7045	GL004 protein	8.70
	433741	AA609019	Hs.159343	ESTs	8.70
	426340	Z97989	Hs.169370	FYN oncogene related to SRC, FGR, YES	8.69
		AA317036 ·	Hs.41989	ESTs	8.67
60		Al225235		Homo sapiens cDNA: FLJ23231 fis, clone C	8.67
	420144	AA811813	Hs.119421		8.66
		AA256756	Hs.31178	ESTs	8.66
		NM_002104	Hs.3066	granzyme K (serine protease, granzyme 3;	8.66
		BE244076	Hs.159578	Homo sapiens mRNA for FLJ00020 protein,	8.65
65		BE246449	Hs.2157	Wiskott-Aldrich syndrome (eczema-thrombo	8.64
		W68180		Homo sapiens cDNA FLJ12507 fls, clone NT	8.64
		AJ001443		splicing factor 3b, subunit 3, 130kD	8.64
		NM_006895	Hs.81182		8.64
		NM_004736		xenotropic and polytropic retrovirus rec	8.63
				-81	

		DE040040	11. 440700	to the the tender of the tender	0.60
	-	BE218919		hypothetical protein FLJ10688	8.63
		AW795196		ring finger protein 14	8.63
		BE242803	Hs.262823	hypothetical protein FLJ10326	8.62
	401851				8.62
5	401866				8.62
		AW996872		a disintegrin and metalloproteinase doma	8.62 8.62
		AA251594		PIBF1 gene product	•
		AW408530		ClpX (caseinolytic protease X, E. coli)	8.62
10		BE550182		RaiGEF-like protein 3, mouse homolog	8.62 8.62
10		Al831594	HS.6864/	ESTs, Weakly similar to ALU7_HUMAN ALU S	8.60
		AW749617	11- 00000	gb:RC3-BT0502-130100-012-g07 BT0502 Homo	8.60
		Al767756	Hs.82302		8.60
		AA829402	Hs.47939		8.60
15		AI972094		Homo sapiens cDNA FLJ13741 fis, clone PL KIAA1634 protein	8.60
13		Al692181 AF009746		ATP-binding cassette, sub-family D (ALD)	8.60
		X54136		immunoglobulin lambda locus	8.60
		U91939		G protein-coupled receptor 25	8.60
	405074	001000	1,0.2.40120	a promin overproving	8.59
20		Al479773	Hs.181679	ESTs	8.59
20		BE328882		ESTs, Moderately similar to U119_HUMAN U	8.58
		AA091228		gb:cchn2152.seq.F Human fetal heart, Lam	8.57
		BE379749	Hs.85201	C-type (calcium dependent, carbohydrate-	8.56
		AL008637		neutrophil cytosolic factor 4 (40kD)	8.56
25		AW947164	Hs.195641		8.56
		AW204272	Hs.199371	ESTs	8.55
	431180	H55883		gb:yq94h03.r1 Soares fetal liver spleen	8.54
	445988	BE007663	Hs.13503	Inactivation escape 2	8.54
	405876				8.54
30	407235	D20569	Hs.169407	SAC2 (suppressor of actin mutations 2, y	8.54
		Al738616		hydroxyprostaglandin dehydrogenase 15-(N	8.54
		AF193612		tunatic fringe (Drosophila) homolog	8.54
		AW082633	Hs.212715		8.54
۰.		AA446183	Hs.91885		8.53
35		Al955765	Hs.146907		8.52
		M31899	Hs.77929	excision repair cross-complementing rode	8.51 8.51
	405552	4141074455	U= 000000	ECTa Maakh similar ta prohil A-hydrovy	8.50
		AW971155	Hs.14373	ESTs, Weakly similar to prolyl 4-hydroxy	8.50
40		AA426117 R68857	Hs.265499		8.50
40		Al765890		ESTs; Moderately similar to IIII ALU SUB	8.50
		AV659397	Hs.282948	· · · · · · · · · · · · · · · · · · ·	8.50
		AW891873	110.202040	gb:CM3-NT0090-040500-173-b02 NT0090 Homo	8.50
		Al927288	Hs.196779		8.48
45		AA279098	Hs.187636		8.48
		AW137635	Hs.44238	ESTs	8.48
	450624	AL043983	Hs.125063	Homo sapiens cDNA FLJ13825 fis, clone TH	8.48
	452573	Al907957	Hs.287622	Homo sapiens cDNA FLJ14082 fis, clone HE	8.48
	456341	AA229126	Hs.122647	N-myristoyitransferase 2	8.48
50		AA593731	Hs.75613	CD36 antigen (collagen type I receptor,	8.47
	446985	AL038704	Hs.156827	ESTs, Weakly similar to ALU1_HUMAN ALU S	8.46
		AL080276	Hs.268562	regulator of G-protein signalling 17	8.46
	400268				8.46
		AW891965	Hs.289109	dimethylarginine dimethylaminohydrolase	8.45
55		NM_014737		Ras association (RaiGDS/AF-6) domain fam	8.44 8.44
		AW057782	Hs.293053		8.44
		AW732240	Hs.300615	gb:wj71c12.x1 NCI_CGAP_Lu19 Homo sapiens	8.42
		Al814302		gb.w// 1012.x1 NOI_OGAF_Lu19 Hollo sapiells	8.42
60	406006	AW971239	Hs.293982	FCTe	8.42
UU		AB014540		SWAP-70 protein	8.42
		AB014540 AW136083		ESTs, Weakly similar to S59501 interfero	8.42
		AW 130063 AI458653	Hs.201881		8.41
		AA358015		gb:EST66864 Fetal lung III Homo sapiens	8.40
65		AW978439	Hs.69504	ESTs	8.40
55		AA013051	Hs.91417	topoisomerase (DNA) II binding protein	8.40
		AW972830		gb:EST384925 MAGE resequences, MAGL Homo	8.40
		AA305688	Hs.267695	UDP-Gal:betaGlcNAc beta 1,3-galactosyltr	8.40
		Al521310	Hs.283365	ESTs, Weakly similar to ALU5_HUMAN ALU S	8.40

		AL122043	Hs.19221	hypothetical protein DKFZp566G1424	8.40
		AW844498	HS.289052	Homo saplens LENG8 mRNA, variant C, part	8.38 8.37
	400817	BE245297		qb:TCBAP1E2482 Pediatric pre-B cell acut	8.37
5	_	AA071267		gb:zm61g01.r1 Stratagene fibroblast (937	8.36
9		BE047779	Hs.44701	ESTs	8.36
		X04430	Hs.93913	interleukin 6 (interferon, beta 2)	8.36
	408327	AW182309	Hs.249963	ESTs, Highly similar to dJ1170K4.4 [H.sa	8.35
	403976				8.34
10		AA379036		gb:EST91809 Synovlal sarcoma Homo sapien	8.33
		AW188551	Hs.99519	Homo sapiens cDNA FLJ14007 fis, clone Y7	8.33
		AW997704 AF119847	Hs.11493	Homo sapiens cDNA FLJ13536 fis, clone PL Homo sapiens PRO1550 mRNA, partial cds	8.32 8.32
		AW937670	Hs.254379	·	8.32
15		NM_015698		T54 protein	8.30
		T70298		gb:yd26g02.s1 Soares fetal liver spleen	8.30
	416852	AF283776	Hs.80285	Homo sapiens mRNA; cDNA DKFZp586C1723 (f	8.30
		AF084866		gb:Homo sapiens envelope protein RIC-3 (8.30
20		Al732694	Hs.98520	ESTs	8.29
20		AW194962	Hs.199028		8.29 8.29
	450783	BE266695		gb:601190242F1 NIH_MGC_7 Homo sapiens cD	8.28
		AF054839	Hs 122540	tetraspan 2	8.28
		AA037675	Hs.152675		8.28
25		AA744488		ESTs, Moderately similar to ALU1_HUMAN A	8.28
	443907	AU076484	Hs.9963		8.27
		AF106069		ubiquitin specific protease 15	8.26
		AA151730		ESTs, Weakly similar to similar to C.ele	8.26
30		AB007918	Hs.169182 Hs.18672	KIAA0449 protein	8.25 8.25
30		AA974411 AW958264		ESTS, Weakly similar to TRHY_HUMAN TRICH	8.24
		AI963740	Hs.46826		8.24
		AW020782	Hs.79881		8.24
	424534	D87682		KIAA0241 protein	8.24
35		U63830		TRAF family member-associated NFKB activ	8.24
		BE263710	Hs.279904		8.22
		Al914699	Hs.13297	ESTs eukaryotic translation initiation factor	8.22 8.22
		BE396283 BE222450	Hs.266390		8.21
40		AA330214	110220000	gb:EST33935 Embryo, 12 week Il Homo sapi	8.21
		AA888624	Hs.19121	adaptor-related protein complex 2, alpha	8.20
٠	416051	AA835868	Hs.25253	Homo sapiens cDNA: FLJ20935 fis, clone A	8.20
		R40739	Hs.21326	ESTs	8.20
45		W25760	Hs.77631	glycine cleavage system protein H (amino	8.20 8.20
43		AU077143 AV654978	Hs.19904	minichromosome maintenance deficient (S. cystathionase (cystathionine gamma-lyase	8.19
		AI828596	Hs.250691		8.18
		NM_015896		BLu protein	8.18
		J04088		topoisomerase (DNA) II alpha (170kD)	8.18
50		M21305		Human alpha satellite and satellite 3 ju	8.18
		U42349		Putative prostate cancer tumor suppresso	8.18
		NM_001838	Hs.1652	chemokine (C-C motif) receptor 7	8.18
		AL133017	Hs.2210	thyroid hormone receptor interactor 3	- 8.17 8.17
55	401519 447188	H65423	Hs.17631	Homo sapiens cDNA FLJ20118 fis. clone CO	8.16
55		Al263293	Hs.152096		8.16
		AJ278120	Hs.4996	DKFZP564D166 protein	8.14
	448556	AW885606	Hs.5064	ESTs	8.14
60		AA278536	Hs.23262	ribonuclease, RNase A family, k6	8.14
60		AI139058	Hs.23296	ESTS	8.14 8.14
		Al018406	Hs.131284 Hs.89271	ESTS ESTS	8.14 8.14
	421129	BE439899	115.032/	LUIS	0.14

TABLE 9A shows the accession numbers for those primekeys lacking a unigeneID in Table 9. For each probeset we have listed the gene cluster number from which the oligonucleotides were designed. Gene clusters were compiled using sequences derived from Genbank ESTs and mRNAs. These sequences were clustered based on sequence similarity using Clustering and Alignment Tools (DoubleTwist, Oakland California). The Genbank accession numbers for sequences comprising each cluster are listed in the "Accession" column.

Unique Eos probeset identifier number

5

10

Pkey: CAT number: Gene cluster number Accession: Genbank accession numbers 15 **CAT** number Accession Pkey 408057 1035720_-1 AW139565 408069 103655_1 H81795 Z42291 R20973 AA046920 408182 104479 1 AA047854 AA057506 AA053841 20 408338 1052148_1 AW867079 AW867086 AW182772 408828 108463_1 BE540279 AW410659 AA057857 R77693 BE278674 AA063426 AW962323 AW408063 AA063503 AA772927 AW753492 BE175371 AA311147 409126 110159_1 409292 111586_1 AA071051 AA070584 AA069938 AA102136 AA074430 AA070266 AA084967 AA126998 409314 111841_1 25 AA071267 T65940 T64515 AA071334 409385 112523 1 AW386461 AW876408 AW386672 AW386599 AW876258 AW386619 AW386289 AW876136 AW876203 AW876213 AW876301 409398 1126716_1 AW876295 AW876349 AW876365 AW876160 AW876369 AW876352 AW876271 AA076769 AA076781 Al087968 409671 114731_1 409768 1154035_1 AW499566 AW502378 AW499522 AW502046 AW502671 AW501917 AW501868 AW501721 AW502813 30 AW502139 AW502432 AW502235 AW501683 AW502647 409841 1156088_1 AW501756 AW502096 AW502465 AW501715 409842 1156119_1 409853 1156226_1 AW502327 AW502488 AW501829 AW502625 AW502687 AW752953 H88044 BE156092 410531 1207200_1 410688 1216101_1 AW796342 AW796356 BE161430 35 AW807057 AW807054 AW807189 AW807193 AW807369 AW807429 AW807364 AW807365 AW807078 AW807256 AW807180 410846 1223902_1 AW807331 AW809637 AW809697 AW810554 AW809707 AW809885 AW810000 AW810088 AW809742 AW809816 AW809749 AW809639 410896 1226053_1 AW809722 AW809836 AW809774 AW810023 AW810013 AW809813 AW809660 AW809728 AW809768 AW809951 AW809657 AW809954 40 411079 123128_1 AA091228 H71860 H71073 AW845985 AW845991 AW845962 411424 1245497_1 AW849292 AW849431 AW849422 AW849428 AW849420 AW849424 AW849427 411499 1248105_1 411507 1248607_1 AW850140 AW850195 AW850192 411534 1248827_1 AW850473 AW850471 AW850431 AW850523 BE074959 AW880160 45 411972 1268491_1 AW893569 AW893571 AW893588 AW893593 412110 1277844_1 412226 1284289 1 W26786 AW998612 AW902272 412257 1285376_1 AW903830 BE071916 AW948126 AW948139 AW948196 AW948145 AW948162 AW948134 AW948127 AW948124 AW948153 AW948157 AW948125 412405 1293012_1 50 AW948131 AW948158 AW948164 AW948151 413260 1356003 1 BE075281 BE075219 BE075123 BE075119 BE075046 BE142098 BE142092 413471 1371778_1 413729 1385114_1 BE159999 BE160056 BE160107 BE160139 414182 142409_1 AA136301 Al381776 AA136321 55 T81668 C19040 C17569 414989 1511339_1 415354 1534763_1 F06495 R24336 R13046 416011 1566439_1 H14487 R50911 Z43216 416475 1596398_1 T70298 H58072 R02750 417380 1672461 1 T06809 N75735 60 419392 1843934_-1 W28573 AW749617 R64714 AA244138 AA244137 BE094019 419541 185724_1 Al909154 AA526337 AA244193 Al909153 419544 185760_2 420819 196721_1 AA280700 AW975494 AA687385 AA285363 AA285333 AA285359 AA285326 AA285350 421245 200620_1 65 N59027 AA314694 N53937 R08100 422673 219674_1

	422695	219996_1	AA315158 AW961298 N76067 AW802759 Al858495 W04474
	422858	222209_1	R35398 BE252178 AA318153
	422940	223106_1	BE077458 AA337277 AA319285
	423730	231462_1	AA330214 AW962519 T54709
5	423790	232031_1	BE152393 AA330984 BE073904
		238731_1	AA339666 AW952809 AA349119
	424606	241409_1	AA343936 AA344060 AW963081
	425265	249175_1	BE245297 AA353976 AW505023
	426959	2738301	
10	430676	32168_1	AF084866 AF084870 AF084864 AF084867 AF084869 AF084865 AF084868 AW818206 AW812038 BE144813 BE144812
			AW812041 AW812040 AW812067 BE061583 BE061604 T05808 AI352469 AA580921 BE141783 BE141782 BE061601
			AW814393 AW885029
		326269_1	AW972830 AA527647 AA489820 AA570362
1.5		328906_1	H55883 AW971249 AA493900 H55788
15		341283_1	H28383 AW972670 H28359 AA525808
		38937_1	T59538 T59589 T59542 AF147374
		41842_1	AJ132085 Z83805
		43393_1	AL050072 AW900148 BE177778 BE177779 AL390180 AA359908
20		43765_1 46858_1	H66948 AF085954 H66949
20		46879_1	H56389 AF085977 H56173
		48675_1	BE410734 BE560117 BE270054 BE296330 BE267957 Al003007 BE545259
		52842_1	AW891873 AW891897 BE564764
		645767_1	Al245701 BE272724
25		71288_1	BE617135 AW504051 AW504283
		74761_1	AA379036 AA150589 Al696854 BE621316
		84655_1	BE266695 BE265474 N53200 BE267333
		85673_1	AA215672 Al696628 AA013335 H86334 AA017006
		921802_1	A1907039 A1907081
30	452560	922216_1	BE077084 AW139963 AW863127 AW806209 AW806204 AW806205 AW806206 AW806211 AW806212 AW806207 AW806208
			AW806210 Al907497
		928309_1	AW838616 AW838660 BE144343 AI914520 AW888910 BE184854 BE184784
		980026_1	U83527 AL120938 U83522
35		1007366_1	AW860158 AW862385 AW860159 AW862386 AW862341 AW821869 AW821893 AW062660 AW062656
33		1224342_1	AW807530 AW807540 AW807537 AW846086 BE141634 AW846089 AW807499 AW807533 AW838499
		1234759_1 1247099_1	BE071874 BE071882 AW820782 AW821007 AW848032 AW848630 AW848478 AW848623 AW848484 AW848169 AW848830 AW848149 AW848119 AW848893 AW848903
	454977	1247099_1	AW848407
	455121	1254674_1	AW857913 AW857916 AW857914 AW861627 AW861626 AW861624
40		1259023_1	AW984111 AW863918 AW663856
		1266449 1	AW877015 AW877133 AW876978 AW877071 AW876988 AW877069 AW877063 AW877013
			AW903533 AW903516 AW903562 BE085202 BE085215 BE085214 BE085209 BE085172 BE085175 BE085193 BE085211
			BE085199
	455982	1396849_1	BE176862 BE176876 BE176947 BE176878
45	456011	1410860_1	BE243628 BE246081 BE247016 BE241984 BE241534 BE246091 BE245679 BE243620 BE245998 BE242329 BE241417
			BE241457 BE242522 BE241989 BE241464
	456023	1416335_1	R00028 BE247630
		360505_1	AW062439 AW751554 AA579463
50		3642251	AA584854
50		399422_1	Al908236 AA663731
		883688_1	A1814302 A1814428
		889426_1	W07808 AI822066
		918957_1	AI903354 AI903489 AI903488 BEG62300 BEG63346 AI006007
55		921149_1 9452401	BE063380 BE063346 Al906097 Al940425
JJ	459234	54024U_•1	COPUPE

TABLE 9B shows the genomic positioning for those primekeys lacking unigene ID's and accession numbers in Table 9. For each predicted exon, we have listed the genomic sequence source used for prediction. Nucleotide locations of each predicted exon are also listed.

Unique number corresponding to an Eos probeset

4693-4857

217175-217446 78627-79583

134394-134812

19197-19502 22733-23007

24755-24969

48154-48499

77842-77954

45096-45229 143025-143467

82849-83627

134445-134750

153027-153262

44340-44559,44790-45059 137113-137814

403259 7770585

403683 7331517

403690 7387384

403708 5705981 403838 4176355

403851 7708872 403976 7657840

404407 7329316

404426 7407959

404632 9796668

404741 8574139

404756 7706327

404946 7382189

405074 7770440

405125 8247873 405172 9966752

55

60

65

Plus

Plus

Minus

Minus

Plus

Plus

Pius

Plus

Plus

Plus

Plus

Plus

Plus

Plus

Plus

Minus

5

Pkey:

	Ref:	Ref:		Sequence source. The 7 digit numbers in this column are Genbank Identifier (GI) numbers. "Dunham I. et al." refers to the publication entitled "The DNA sequence of human chromosome 22." Dunham I. et al., Nature (1999) 402:489-495.					
10	Strand: Nt_posit	Strand: Nt_position:		Indicates DNA strand from which exons were predicted. Indicates nucleotide positions of predicted exons.					
	Pkey	Ref	Strand	Nt_position					
15	400452	8113550	Minus	90308-90505					
		9801261	Plus	208453-208528,209633-209813					
		9908994	Plus	118036-118166,118681-118807					
		8567867	Minus	174571-174856					
		8569994	Plus	170793-170948					
20		9931121	Plus	29235-29336,36363-36580	•				
20		9958187	Minus	58242-58733					
		7651921	Minus	52033-52158,53956-54120,54957-55052,55420-55480,56452-56666,57221-57718					
		7658481	Plus	192667-192826,194387-194876					
		8096825	Plus	159197-159320					
25		8117619	Plus	73501-73674					
		8570296	Minus	124181-124391					
		6981820	Plus	5302-5545	•				
		9743387	Minus	138534-138629,139234-139294,140121-140335,142033-142479					
		9801342	Minus	147036-147318					
30		6850939	Minus	58360-58545					
50		6433826	Plus	13056-13482					
		7630851	Plus	29929-30126					
		6649315	Plus	157315-157950					
		9838136	Plus	128526-128704,130755-130860					
35		2576349	Plus	14595-14751					
-		7770425	Minus	146443-146664,147794-147971,148351-148480,148980-149111,149801-149949					
		8018106	Plus	73126-73623					
		7690131	Plus	104382-104527,106136-106372	· ·-i				
		9211204	Minus	40403-41961	•1				
40		9908890	Minus	174893-175050,183210-183435					
		9796102	Plus	98273-101430					
		3287156	Minus	53242-53432					
		6010110	Plus	25026-25091,25844-25920					
		8918414	Plus	69071-69642					
45		9187337	Plus	26961-27101					
		9369121	Minus	32589-32735,35478-35666	•				
		9369121	Minus	76355-76479					
•		9967547	Plus	85537-85671,86379-86469					
		9581599	Minus	46624-46784					
50		9211494	Minus	92349-92572,92958-93084,93579-93712,93949-94072,94591-94748,95214-95337					
		7637807	Plus	7271-7527					
		7770505	Dire	4603.4957					

	405236 7249076	Minus	151699-151915
	405325 6094661	Minus	25818-26380
	405411 3451356	Minus	17503-17778,18021-18290
	405495 8050952	Minus	72182-72373
5	405552 1552506	Plus	45199-45647
•	405601 5815493	Minus	147835-147935,149220-149299
	405685 4508129	Minus	37956-38097
	405777 7263187	Minus	104773-105051
	405856 7653009	Plus	101777-102043
10	405876 6758747	Plus	39694-40031
	405932 7767812	Minus	123525-123713
	405934 6758795	Plus	159913-160605
	406006 8247801	Minus	42640-42776
	406134 9163473	Plus	153291-153452
15	406189 7289992	Minus	22007-22234
	406422 9256411	Plus	163003-163311
	406516 7711422	Minus	128375-128449.128560-128784
	406538 7711478	Plus	35196-35367,38229-38476,40080-40216,43522-43840
	406554 7711566	Plus	106956-107121
20	406577 7711730	Plus	11377-11509

TABLE 10: shows genes, including expression sequence tags differentially expressed in taxol resistant prostate tumor xenografts as compared to taxol sensitive prostate tumor
 xenografts. The genes are indicated as either being upregulated or downregulated during the induction of taxol resistance in sequential passages of the grafts.

10 15	ExAccn: Exemplar of UnigenelD: Unigene no Unigene Title: Unigene g Eos: Internal Ec		Exemplar A Unigene nu Unigene ge Internal Eos	Inique Eos probeset identifier number Exemptar Accession number, Genbank accession number Inigene number Inigene gene title nternal Eos name assage number																
13																				
	Pkey	ExAcon	UnigenelD	UnigenTitle	Eos	Resp.	F00	F00	F02	F02	F05	F05	F07	F09	F10	F11	F13	F14		
20	117921	N51002	Hs.47170	Liprin A2	PM28	UP	1	9	8	9	32	20	34	122	105	82	71	111		
	112971		Hs.4299	ESTs	CHA1	down	290	281	267	335	270	284	150	157	83	89	49	75		
		Al167942	Hs.61635	STEAP	PAA5	down	106	111	103	71	34	67	33	14	2	1	1	1		
	119018	N95796	Hs.179809	ESTs	PAB2	down	765	841	757	909	742	704	478	428	253	175	228	238		
	110844	N31952	Hs.167531	ESTs	PAV7	down	175	192	147	141	123	129	73	65	55	48	54	84		
25	100654	HG2841-HT29	169	Hs.75442	Album	in, A	PM01	down	666	605	504	728	357	445	602	187	117	127	117	113
	100655	HG2841-HT29	170	Hs.75442	Album	in, A	PM02	down	620	653	486	688	368	386	606	175	101	95	115	97
	102076	U09579	Hs.252437	cyclin-dep	PM03	down	101	94	143	190	105	107	88	40	34	31	46	22		
	102208	U22961	Hs.75442	albumin	PM04	down	495	424	323	518	252	296	467	188	169	143	165	145		
	103739	AA075779	-	mitochondr	PM05	down	75	190	606	230	378	106	218	88	69	192	69	99		
30	107036	AA599690'	Hs.15725	SBB148	PM06	down	87	124	115	188	132	111	66	71	49	70	38	50		
	108242	AA062746	-	ESTs	PM07	down	14	20	252	13	22	43	193	10	10	104	21	18		
	108282	AA065143	-	solute car	PM08	down	27	54	178	73	108	37	53	24	14	53	15	34		
	108679	AA115963	-	beta-1-glo	PM09	down	680	893	1292	656	869	389	1	74	118	662	359	409		
	108731	AA126313	Hs.107476	ATP syntha				19	185	25	60	1	32	3	7	14	1	1	•	
35	110675	H89355	Hs.6598	adrenergic	PM11	down	207		237	239	231	220	119	145	93	64	56	124		
		AA283804	Hs.193552		PM12				282	271	340	334	115	238	100	196	83	207		
		AA430124			PM13			93	94	154	132	91	23	54	23	76	14	41		
		AA281591			PM14			157	58	141	159	127	39	83	35	37	16	46		
		Y00705	Hs.181286		PM15				214	150	106	128	177	85	54	63	66	56		
40		AA490775	Hs.5920	N-ecetylma				150	132	178	126	139	53	94	48	67	41	80		
		AA032221			PM17					215	205	180	132	65	68	50	48	63		
		AA283085	Hs.64065		PM18			148	161	150	92	108	42	99	42	65	29	126		
		D62633	Hs.8236		PM19			240		212	231	189	89	123	107	95	68	91		
	135400	M23263	Hs.99915	androgen r	PM20	down	36	167	99	178	132	101	23	71	26	122	14	44		
45																				

TABLE 11: shows genes, including expression sequence tags that are up-regulated in prostate tumor tissue compared to normal prostate tissue as analyzed using Affymetrix/Eos Hu01 GeneChip array. Shown are the ratios of "average" normal prostate to "average" prostate cancer tissues.

	Pkey: ExAccn: UnigenelD: Unigene Title:	Exemplar Accessi Unigene number Unigene gene title			·
	R1:	Background subtr	acted normal prostate :	prostate tumor tissue	
	Pkey	ExAcon	UnigenelD	Unigene Title	R1
	101336	L49169	Hs.75678	FBJ murine osteosarcoma viral oncogene homolog B	0.012
10	130642	M63438	Hs.156110	Immunoglobulin kappa variable 1D-8	0.015
	133512	X01677	Hs.195188	glyceraldehyde-3-phosphate dehydrogenase	0.017
	133436	H44631	Hs.737	immediate early protein	0.017
	129292	X13810	Hs.1101	POU domain; class 2; transcription factor 2	0.019
	100610	HG2566-HT4792	•	Microtubule-Associated Protein Tau, Alt. Splice 3, Exon 8	0.02
15	133448	M34516	Hs.170116	immunoglobulin lambda-like polypeptide 3	0.021
	125193	W67577	Hs.84298	CD74 antigen (invariant polypeptide of major histocompatibility	
				complex; class II antigen-associated)	0.022
	133456	T49257	Hs.183704	ubiquitin C	0.022
	134546	AA459310	Hs.8518	Homo sapiens mRNA; cDNA DKFZp586L1722 (from clone	
20				DKFZp586L1722)	0.023
	102131	U15085	Hs.1162	major histocompatibility complex; class II; DM beta	0.023
	101375	M13560	Hs.84298	CD74 antigen (invariant polypeptide of major histocompatibility	
	•	•		. complex; dass II antigen-associated)	0.023
	100674	HG3033-HT3194		Spliceosomal Protein Sap 62	0.024
25	134365	R32377	Hs.82240	syntaxin 3A	0.027
	132335	D60387	Hs.189885	ESTs	0.027
	110303	H37901	Hs.32706	ESTs ·	0.028
	131678	N59162	Hs.30542	ESTs	0.028
	116599	D80046	Hs.250879	ESTs .	0.029
30	133769	M17733	Hs.75968	thymosin; beta 4; X chromosome	0.029
	107904	AA026648	Hs.61389	ESTs	0.03
	129427	T80746	Hs.111334	ferritin; light polypeptide	0.03
	105987	AA406631	Hs.110299	mitogen-activated protein kinase kinase 7	0.03
	131466	F03233	Hs.27189	ESTs	0.032
35	102859	X00274	Hs.76807	Human HLA-DR alpha-chain mRNA	0.032
	134626	S82198	Hs.8709	caldecrin (serum calcium decreasing factor; elastase IV)	0.032
	134170	M63138	Hs.79572	cathepsin D (lysosomal aspartyl protease)	0.033
	131713	X57809	Hs.181125	immunoglobulin tambda gene cluster	0.034
	100748	HG3517-HT3711		Alpha-1-Antitrypsin, 5' End	0.034
40	118769	N74496		ESTs .	0.034
	111734	R25375	Hs.126916	ESTs	0.036
	109221	AA192755	Hs.85840	ESTs; Weakly similar to stac [H.sapiens]	0.036
	133846	AA480073	Hs.76719	U6 snRNA-associated Sm-like protein	0.036
	135281	AA401575	Hs.97757	ESTs -	0.037
45	119073	R32894	Hs.45514	v-ets avian erythroblastosis virus E26 oncogene related	0.037
	100760	HG3576-HT3779		Major Histocompatibility Complex, Class II Beta W52	0.037
	101426	M19483 .	Hs.25	ATP synthase; H+ transpring; mitochndrl F1 complex; beta polypept	0.038
	129568	AA428025	Hs.114360	transforming growth factor beta-stimulated protein TSC-22	0.038
~~	130900	Z38468	Hs.21036	ESTs; Moderately similar to F25965_3 [H.sapiens]	0.039
50	133879	M13829	Hs.77183	v-raf murine sarcoma 3611 viral oncogene homolog 1	0.039
	100627	HG2702-HT2798		Serine/Threonine Kinase (Gb:Z25424)	0.039
•	129424	M55593	Hs.111301	matrix metalloproteinase 2 (gelatinase A; 72kD gelatinase; 72kD type IV collagenase)	0.039
	128652	AA621245	Hs.103147	ESTs; Weakly similar to similar to SP:YR40_BACSU [C.elegans]	0.039
55	129979	T72635	Hs.13956	ESTS	0.039
	133468	X03068	Hs.73931	major histocompatibility complex; class II; DQ beta 1	0.04
	102636	U67092		Human ataxia-telangiectasia locus protein (ATM) gene, exons	
				1a, 1b, 2, 3 and 4, partial cds	0.04
	129536	M33493	Hs.184504	tryptase; alpha	0.04
60	133599	M64788	Hs.75151	RAP1; GTPase activating protein 1	0.041
				= *	

	102104	U12139		Human alpha1(XI) collagen (COL11A1) gene, 5' region and exon 1	0.041
	131340	AA478305	Hs.25817	Homo sapiens chromosome 19; cosmid R27216	
					0.041
	130446	X79510	Hs.155693	protein tyrosine phosphatase; non-receptor type 21	0.042
_	101352	L77701	Hs.16297	COX17 (yeast) homolog; cytochrome c oxidase assembly protein	0.042
5	122593	AA453310	Hs.128749	alpha-methylacyl-CoA racemase	0.042
	130181	R39552	Hs.151608	Homo sapiens clone 23622 mRNA sequence	0.042
	134071	Z14093	Hs.78950	branched chain keto add dehydrogenase E1; alpha polypeptide	0.042
	134071	214033	H2.70930		
				(maple syrup urine disease)	0.042
	108129	AA053252	Hs.185848	ESTs; Weakly similar to II ALU SUBFAMILY J WARNING	
10				ENTRY !! [H.sapiens]	0.043
	130511	L32137	Hs.1584	cartilage oligomeric matrix protein (pseudoachondroplasia;	0.0-0
	.00011	202107	110.1004	callinge digomente manis protein (pseudoacijonuju)asia,	
	400000	4.4004.550	11	epiphyseal dysplasia 1; multiple)	0.043
	133336	AA291456	Hs.71190	ESTs	0.043
	132982	L02326	Hs.198118	immunoglobulin lambda-like polypeptide 2	0.044
15	131880	AA047034	Hs.33818	RecQ protein-like 5	0.044
_	130540	U35234	Hs.159534	protein tyrosine phosphatase; receptor type; S	0.044
	133467	AA258595			
			Hs.73931	major histocompatibility complex; class II; DQ beta 1	0.044
	101191	L20688	Hs.83656	Rho GDP dissociation inhibitor (GDI) beta	0.044
	101860	M95610	Hs.37165	collagen; type iX; alpha 2	0.044
20	102799	U88898		Human endogenous retroviral H protease/integrase-derived ORF1	•••
				mRNA, complete cds, and putative envelope prot mRNA, partial cds	0.044
	107000	Dangen	Un FOOD		0.044
	107200	D20350	Hs.5628	ESTs	0.044
	101166	L14927	Hs.2099	lipocalin 1 (protein migrating faster than albumin; tear prealbumin)	0.044
	134289	M54915	Hs.81170	pim-1 oncogene	0.044
25	135329	AA436026	Hs.98858	ESTs	0.044
	124950	T03786	Hs.151531	protein phosphatase 3 (formerly 2B); catalytic subunit; beta isoform	0.044
	12-1000	100100	110.101001		
				(calcineurin A beta)	0.044
	102919	X12447	Hs.183760	aldolase A; fructose-bisphosphate	0.044
	100574	HG2279-HT2375		Triosephosphate Isomerase	0.045
30	131286	AA450092	Hs.25300	Homo sapiens clones 24718 and 24825 mRNA sequence	0.045
	102675	U72512		Human B-cell receptor associated protein (hBAP) alternatively	0.040
	102075	G/ES/E			
				spliced mRNA, partial 3'UTR	0.045
	131332	R50487	Hs.25717	ESTs	0.045
	101634	M57731	Hs.75765	GRO2 oncogene	0.046
35	113118	T47906	Hs.220512	ESTs	0.046
	124884	R77276	Hs.120911	ESTs	
					0.046
	130523	W76097	Hs.214507	ESTs	0.046
	110244	H26742	Hs.25367	ESTs; Weakly similar to ALR [H.sapiens]	0.046
	131932	AA454980	Hs.25601	chromodomain helicase DNA binding protein 3	0.046
40	132509	H09751	Hs.5038	neuropathy target esterase	0.046
	133372	AA291139	Hs.72242	ESTs	0.046
	100817		110.7 ELTE		
		HG4011-HT4804		Dystrophin-Associated Glycoprotein, 50 Kda, Alt. Splice 2	0.047
	106746	AA476436	Hs.7991	ESTs	0.047
	135401	L14813	Hs.169271	carboxyl ester lipase-like (bile salt-stimulated lipase-like)	0.047
45	130479	R44163	Hs.12457	Homo sapiens clone 23770 mRNA sequence	0.047
	102589	U62015	Hs.8867	cysteine-rich; angiogenic inducer; 61	0.047
	121521				
		AA412165	Hs.97358	EST	0.048
	135340	AA425137	Hs.99093	Homo sapiens chromosome 19; cosmid R28379	0.048
	132336	AA342422	Hs.45073	ESTs	0.048
50	115368	AA282133	Hs.88960	ESTs; Weakly similar to similar to collagen [C.elegans]	0.048
	101278	L38487	Hs.110849	estrogen-related receptor alpha	0.048
	103284	X80200	Hs.8375		
			П8.03/Э	TNF receptor-associated factor 4	0.048
	100564	HG2239-HT2324		Potassium Channel Protein (Gb:Z11585)	0.048
	133132	Z40883	Hs.65588	ESTs; Weakly similar to dJ393P12.2 [H.sapiens]	0.048
55	121811	AA424535	Hs.98416	ESTs	0.048
	129613	AA279481	Hs.238831	ESTs; Weakly similar to collagen alpha 1(XVIII) chain [M.musculus]	0.049
	132468	S79854		delodinase; lodothyronine; type [[]	
•			Hs.49322		0.049
	120111	W95841	Hs.136031	ESTs .	0.049
	103668	Z83741	Hs.248174	H2A histone family; member M	0.049
60	130386	F10874	Hs.234249	mitogen-activated protein kinase 8 interacting protein 1	0.049
	104275	C02170	Hs.39387	ESTs; Weakly smir to weak smirity to ribosomal prot L14 [C.elegans]	0.049
	106305	AA436146	Hs.12828	ESTs	0.05
	116431	AA609878	Hs.55289	ESTs; Weakly smir to 110 KD CELL MEMBRANE GLYCOPROTEIN [H.sapiens]	0.813
	120339	AA206465	Hs.256470	EST	0.05
65	114427	AA017063		ESTs; Highly similar to Miz-1 protein [H.sapiens]	0.05
	118821	N79070	Hs.94789	ESTs	0.05
	118979	N93798	Hs.43666	protein tyrosine phosphatase type IVA; member 3	0.05
	107495	W78776	Hs.90375	ESTs	0.051
	120240	Z41732	Hs.66049	ESTs	0.051

	114331	Z41309	Hs.12400	ESTs	0.051
	130947	R40037	Hs.21506	ESTs	0.052
	129242	W81679	Hs.5174	ribosomal protein \$17	0.052
	131413	AA482390	Hs.26510	ESTs; Modly smlr to vacuolar prot sorting homolog r-vps33b [R.norvegicus]	0.052
5	112304	R54798	Hs.26239	ESTs	0.052
	101416	M17254	Hs.45514	v-ets avian erythroblastosis virus E26 oncogene related	0.052
	131201	AA426304	Hs.24174	ESTs	0.052
•	101054	K02405	Hs.73933	Human MHC class II HLA-DQ-beta mRNA (DR7 DQw2); complete cds	0.052
	101306	L41143	Hs.232069	T-cell leukemia translocation altered gene	0.053
10	129311	T55087		yb45c08.r1 Stratagene fetal spleen (#937205) Homo sapiens cDNA	
				done IMAGE:74126 5', mRNA sequence.	0.053
	129942	U95301	Hs.144442	phospholipase A2; group X	0.053
	119210	R93340	Hs.92995	ESTs ·	0.053
	101046	K01160		Accession not listed in Genbank	0.053
15	114086	Z38266	Hs.12770	Homo sapiens PAC clone DJ0777O23 from 7p14-p15	0.053
	110171	H19964	Hs.31709	ESTs	0.053
	101004	J04101	Hs.248109	v-ets avian erythroblastosis virus E26 oncogene homolog 1	0.053
	129715	N58479	Hs.12126	ESTs; Weakly similar to LR8 [H.sapiens]	0.053 0.053
20	101581	M34996	Hs.198253	major histocompatibility complex; class II; DQ alpha 1	0.053
20	113285	T66830	Hs.182712	ESTS	0.054
	127537	AA569531	Hs.162859	ESTs Con Freithad Doo Clope \$10	0.054
	100813	HG3995-HT4265	Un 70000	Cpg-Enriched Dna, Clone S19 3-hydroxybutyrate dehydrogenase (heart; mitocfiondrial)	0.054
	101841	M93107 R77159	Hs.76893	ESTs	0.054
25	135053	M17886	Hs.93678 Hs.177592	ribosomal protein; large; P1	0.054
43	101419 119724	W69468	Hs.47622	ESTs	0.055
	102673	U72509	113.47 022	Human alternatively spliced B8 (B7) mRNA, partial sequence	0.055
	129877	AA248589	Hs.13094	ESTs; Weakly similar to ORF YGR101w [S.cerevisiae]	0.055
	114788	AA156737	Hs.103904	EST	0.055
30	123812	AA620607	Hs.111591	ESTs	0.055
	117669	N39237	Hs.44977	ESTs	0.055
	123782	AA610111	Hs.162695	EST	0.055
	102395	U41767	Hs.92208	a disintegrin and metalloproteinase domain 15 (metargidin)	0.055
	133795	M12529	Hs.169401	apolipoprotein E	0.055
35	123193	AA489228	Hs.136956	ESTs	0.056
	132595	AA253369	Hs.155742	glyoxylate reductase/hydroxypyruvate reductase	0.056
	104161	AA456471	Hs.7724	KIAA0963 protein	0.056 0.056
	115330	AA281145	Hs.88827	ESTs	0.056
40	112893	T08000	Hs.194684	bassoon (presynaptic cytomatrix protein)	0.056
40	133475 128699	L29217 K03207	Hs.73987 Hs.103972	CDC-like kinase 3 proline-rich protein BstNI subfamily 4	0.056
	102940	X13956	Hs.24998	Hu 12S RNA induced by poly(rl); poly(rC) and Newcastle disease virus	0.056
	131299	AA431464	Hs.25426	ESTs; Weakly similar to unknown [H.sapiens]	0.057
	102495	U51240	Hs.79356	Lysosomal-associated multispanning membrane protein-5	0.057
45	129594	R70379	Hs.115396	Human germline IgD chain gene; C-region; C-delta-1 domain	0.057
	118593	N69020	Hs.207689	EST	0.057
	126702	U54602	Hs.2785	keratin 17	0.057
	124386	N27368	Hs.212414	sema domain; immunoglobulin domain (lg); short basic domain;	
				secreted; (semaphorin) 3E	0.057
50	130538	M20786	Hs.159509	alpha-2-plasmin inhibitor	0.057
	114299	Z40782	Hs.22920	similar to \$68401 (cattle) glucose induced gene	0.057
	115604	AA400378	Hs.49391	ESTs	0.057
	106052	AA416947	Hs.6382	ESTs; Highly similar to KIAA0612 protein [H.sapiens]	0.057
	131730	U05681	Hs.31210	B-cell CLL/lymphoma 3	0.057
55	131285	AA479498	Hs.25274	ESTs; Modly smlr to putative seven pass transmembrane prot [H.sapiens]	0.058 0.058
	129705	X78706	Hs.12068	camitine acetyltransferase	0.058
	123175	AA489010 Z30644	Hs.178400 Hs.123059	ESTs chloride channel Kb	0.058
	103592 118196	N59478	Hs.48396	ESTs: Moderately similar to tumor necrosis factor-alpha	0.000
60	110190	1100410	115.40550	-induced protein B12 [H.sapiens]	0.058
00	104886	AA053348	Hs.144626	growth differentiation factor 11	0.058
	104250	AF000575	Hs.105928	leukocyte immunoglobulin-like receptor; subfamily B (with TM	
				and ITIM domains); member 3	0.058
	113301	T67452	Hs.13104	EST	0.058
65	110441	H50302	Hs.19845	ESTs; Highly smlr to prot phosphatase 2A BR gamma subunit [H.saplens]	0.058
	125297	Z39215	Hs.159409	ESTs	0.058
	135258	AA292423	Hs.97272	ESTs; Weakly similar to dJ281H8.2 [H.sapiens]	0.058
	130633	T92363	Hs.178703	ESTs	0.058
	112006	R42607	Hs.22241	hypothetical protein	0.058

	130805	U12194	Hs.170238	sodium channel; voltage-gated; type I; beta polypeptide	0.058
	134907	D80002	Hs.178292	KIAA0180 protein	0.058
	132619	AA404565	Hs.53447	ESTs; Moderately similar to kinesin light chain 1 [M.musculus]	0.058
	135115	N35489	Hs.94653	neurochondrin	0.058
5	100531	HG1872-HT1907		Major Histocompatibility Complex, Dg	0.058
•	124530	N62256	Hs.102727	EST	0.058
	119960	W87533	Hs.32699	ESTs; Moderately similar to LIV-1 protein [H.saplens]	0.058
	132793	AA478999	Hs.56966	KIAA0906 protein	0.058
	101076	L04270	Hs.1116	lymphotoxin beta receptor (TNFR superfamily; member 3	0.058
10	130655	N92934	Hs.17409	cysteine-rich protein 1 (intestinal)	0.058
10	134458	AA192614	Hs.83577	cysteine and glycine-rich protein 3 (cardiac LIM protein)	0.058
	105904	AA401452	Hs.32060	ESTs	0.059
•	132878	AA026793	Hs.58679	ESTs; Weakly similar to 4F2/CD98 light chain (M.musculus)	0.059
	121828	AA425166	Hs.98497	ESTs	0.059
15	133418	U76366	Hs.172727	Treacher Collins-Franceschetti syndrome 1	0.059
13	129317	N46244	Hs.110373	ESTs	0.059
	130153	D85815		ras homolog gene family; member D	0.059
			Hs.15114 Hs.102493	ESTs	0.059
	124403	N31745	Hs.134170	ESTS	0.059
20	127683	AA668123		==::	
20	129814	W20070 D59682	Hs.168625	KIAA0979 protein	0.059
	131770		Hs.31833	ESTs	0.06
	117557	N33920	Hs.44532	diubiquitin	0.06
	103522	Y10514	Lie oroczo	H.sapiens mRNA for CD152 protein	0.06
25	120029	W91960	Hs.250640	sequence-specific single-stranded-DNA-binding protein	0.06
23	102135	U15460	Hs.41691	activating transcription factor B	0.06
	123617	AA609183	Hs.181131 Hs.9739	ESTS	0.06
	112136	R46100		ESTs	0.061
	133725	V00563	Hs.179543	immunoglobulin mu	0.061
30	102069	U09196	Hs.82520	Hu 1.1 kb mRNA upregltd in retinoic acid treated HL-60 neutrophilic cells	0.061
20	106555	AA455000	Hs.16725	ESTs	0.061
	123269	AA491226	Hs.105280	ESTs; Weakly similar to dJ963K23.2 [H.sapiens]	0.061
	109088	AA166837	Hs.72620	DKFZP434I114 protein	0.061
	129399	AA263028	Hs.111076	malate dehydrogenase 2; NAD (mitochondrial)	0.061
35	129375	W79850	Hs.11081	ESTs; Weakly similar to HPBRII-7 protein [H.sapiens]	0.061
33	135271	AA397763	Hs.97562	ESTs	0.061
	132958	W90398	Hs.6147	KIAA1075 protein	0.061
	129364	AA477106	Hs.110757	DNA segment on chromosome 21 (unique) 2056 expressed sequence	0.061
	123427	AA598548	Hs.112471	ESTs	0.061
40	105236	AA219179	Hs.19105	translocase of inner mitochondrial membrane 17 (yeast) homolog B	0.061
40	101012	J04444	Hs.697	cytochrome c-1	0.062
	134791	L18983	Hs.89655	protein tyrosine phosphatase; receptor type; N	0.062
	133700	K01396	Hs.75621	protease inhibitor 1 (anti-elastase); alpha-1-antitrypsin	0.062
	123887	AA621065	Hs.112943	ESTS	0.062
45	129363 105719	H05704	Hs.110746 Hs.36793	H sapiens HCR (a-helix coiled-coil rod homologue) mRNA; complete cds	0.062
43		AA291644		ESTs	0.062
	124226	· H62396	Hs.190266	ESTS	0.062
	117437	N27645		yw5e3.s1 Weizmann Olfactory Epithelium H sapiens cDNA clone	0.000
	100744	A 4004400	Hs.55898	IMAGE:255676 3' smlr to contains L1.13 L1 repetitive element;, mRNA seq	0.062
50	132741 134437	AA394133	Hs.198253	ESTs; Highly similar to OASIS protein [M.musculus]	0.062 0.062
50		M26041		major histocompatibility complex; class II; DQ alpha 1	
	107664	AA010594	Hs.5326	ESTs; Moderately similar to pim-1 protein [H.saptens]	0.062
	120844	AA349417	Hs.96917	ESTs	0.062
	101574	M34182	Hs.158029	protein kinase; cAMP-dependent; catalytic; gamma	0.062
55	131219	C00476	Hs.24395	small inducible cytokine subfamily B (Cys-X-Cys); member 14 (BRAK)	0.062
23	103495	Y09022	Hs.153591	Not56 (D. melanogaster)-like protein	0.062
	129607	AA404594	Hs.11607	ESTs	0.062
	106467	AA450040	Hs.154162	ADP-ribosylation factor-like 2	0.062
	128841	T16358	Hs.106443	ESTs	0.062
60	100515	HG1723-HT1729		Macrophage Scavenger Receptor, Alt. Splice 2	0.062
60	119332	T54095	U- 00440	ESTs; Weakly similar to !! ALU SUBFAMILY J WARNING ENTRY !! [H.sapiens]	0.062
	134516	AA171939	Hs.23413	ESTs	0.062
	135012	X73608	Hs.93029	sparc/osteonectin; cwcv and kazal-like domains proteoglycan (testican)	0.063
	103575	Z26256	11 88455	H.sapiens isoform 1 gene for L-type calcium channel, exon 1	0.063
C F	115514	AA297739	Hs.55609	ESTs; Weakly similar to ISOLEUCYL-TRNA SYNTHETASE;	
65		44004077		CYTOPLASMIC [H.sapiens]	0.063
	103996	AA321355	11	EST2393 Bone marrow Homo sapiens cDNA 5' end, mRNA sequence	0.063
	110505	H55992	Hs.20495	DKFZP434F011 protein	0.063
	133912	X62744	Hs.77522	major histocompatibility complex; class II; DM alpha	0.063
	129581	M33600	Hs.180255	major histocompatibility complex; class II; DR beta 1	0.063

	130139	R38280	Hs.150922	BCS1 (yeast homolog)-like	0.064
	105817	AA397825	Hs.5307	synaptopodin	0.064
	134658	AA410617	Hs.178009	ESTs	0.064
_	100306	D50495	Hs.80598	transcription elongation factor A (SII); 2	0.064
5	100277	D42053	Hs.75890	site-1 protease (subtilisin-like; sterol-regulated; cleaves sterol regulatory	
_				element binding proteins)	0.064
	133116	D61259	Hs.6529	ESTs	0.064
	134909	AA521488	Hs.90998	KIAA0128 protein	0.064
	130319	X74794	Hs.154443	minichromosome maintenance deficient (S. cerevisiae) 4	0.064
10	132057	AA102489	Hs.173484	ESTs	0.064
	108334	AA070473		zm7c8.s1 Stratagene neuroepithelium (#937231) Homo sapiens cDNA	
	100007	701070410			0.064
		*	11 10070	clone IMAGE:5399 3', mRNA sequence	
	129763	F10815	Hs.12373	KIAA0422 protein	0.064
	135112	T67464	Hs.94617	ESTs; Weakly similar to predicted using Genefinder [C.elegans]	0.064
15	122269	AA436856	Hs.98910	ESTs	0.064
	133082	AA457129	Hs.6455	RuvB (E coli homolog)-like 2	0.064
	113213	T58607	110.0400	ya94a02.s1 Stratagene placenta (#937225) Homo saplens cDNA clone	0.001
	113213	130007			0.005
				IMAGE:69290 3', mRNA sequence.	0.065
	106228	AA429290	Hs.17719	ESTs	0.065
20	130192	Y12661	Hs.171014	VGF nerve growth factor inducible	0.065
	104894	AA054087	Hs.18858	phospholipase A2; group IVC (cytosolic; calcium-independent)	0.065
	103508	Y10141	12.10000	H.saplens DAT1 gene, partial, VNTR	0.065
			11- 400000		
	128474	U40671	Hs.100299	igase III, DIVA, ATT-departerit	0.065
	134012	AA417821	Hs.237924	ESTs; Highly similar to CGI-69 protein [H.saplens]	0.065
25	134536	AA457735	Hs.850	IMP (inosine monophosphate) dehydrogenase 1	0.065
	111714	R23146	Hs.23466	ESTs	0.065
	110521	H57060	Hs.108268	ESTs	0.065
					0.065
•	103282	X80198	Hs.77628	steroidogenic acute regulatory protein related	
	113921	W80730	Hs.28355	ESTs	0.065
30	129331	N93465	Hs.110453	ESTs; Highly similar to CGI-38 protein [H.saplens]	0.065
	111316	N74597	Hs.180535	ESTs; Weakly similar to mitogen inducible gene mig-2 [H.sapiens]	0.065
	135138	AA036794	Hs.95196	ESTs; Weakly similar to T20B12.3 [C.elegans]	0.065
	107289	T10792	Hs.172098	ESTs .	0.065
25	121405	AA406083	Hs.98007	ESTs	0.065
35	124965	T16275	Hs.106359	ESTs	0.065
	106595	AA456933	Hs.174481	ESTs	0.066
	100106	AF015910		Homo sapiens unknown protein mRNA, partial cds	0.066
	134715	AA282757	Hs.89040	prepronociceptin	0.066
				TVDO	0.066
40	135367	AA480109	Hs.9963	TYRO protein tyrosine kinase binding protein	
40	111533	R08548	Hs.251651	EST	0.066
	128509	R53109	. Hs.247362	dimethylarginine dimethylaminohydrolase 2	0.066
	101030	J05037	Hs.76751	serine dehydratase	0.066
	102753	U80226		Human gamma-aminobutyric acid transaminase mRNA, partial cds	0.067
		R31652	Hs.821	bigiyean	0.067
45	126991				
43	109583	F02322	Hs.26135	ESTs	0.067
	119241	T12559	Hs.221382	ESTs	0.067
	130569	AA156597	Hs.256441	EST; Moderately similar to CGI-136 protein [H.sapiens]	0.067
	112926	T10316	Hs.4302	ESTs	0.067
	120495	AA256073	Hs.190626	ESTs	0.067
50	130931	AA278412	Hs.21346	ESTs; Weakly similar to F42C5.7 gene product [C.elegans]	0.067
50					
	129982	M87789	Hs.140	immunoglobulin gamma 3 (Gm marker)	0.067
	133832	H03387	Hs.241305	estrogen-responsive B box protein	0.067
	110697	H93721	Hs.20798	ESTs -	0.067
	121183	AA400138	Hs.97703	ESTs	0.067
55				Wiskott-Aldrich syndrome (ecezema-thrombocytopenia)	0.067
JJ	130953	U12707	Hs.2157		
	102218	U24183	Hs.75160	phosphofructokinase; muscle	0.067
	114181	Z39079	Hs.8021	KIAA1058 protein	0.067
	116581	D51287	Hs.82148	ribosomal protein S12	0.067
	132498	T87708	Hs.50098	ESTs	0.068
60	103788	AA096014	Hs.9527	ESTs; Highly similar to HSPC013 [H.saplens]	0.068
00			110.0321	Human amiloride-sensitive epithelial sodium channel gamma subunit mRNA,	0.000
	102459	U48936			0.000
				5' end, partial cds	0.068
	100373	D79999	Hs.77225	ADP-ribosyltransferase (NAD+; poly (ADP-ribose) polymerase)-like 1	0.068
	132717	AA203321	Hs.151696	DKFZP727G051 protein	0.068
65	128863	D87462	Hs.106674	BRCA1 associated protein-1 (ubiquitin carboxy-terminal hydrolase)	0.068
				ESTs	0.068
	115193	AA262029	Hs.88218		
	124558	N66046	Hs.141605	ESTs	0.069
	117225	N20392	Hs.42846	ESTs	0.069
	110665	H83380	Hs.32757	ESTs	0.069

				•	
	400005	1 170000	H- 40000F	Variant like feeter A feeth	0.069
	132905	U70663	Hs.182965	Kruppel-like factor 4 (gut)	
	105778	AA348910	Hs.153299	DOM-3 (C. elegans) homolog Z	0.069
	134770	R72079	Hs.89575	CD79B antigen (immunoglobulin-associated beta)	0.069
_	123097	AA485869	Hs.105671	ESTs	0.069
5	100750	HG3523-HT4899		Proto-Oncogene C-Myc, Alt. Splice 3, Orf 114	0.069
	125091	T91518		ye20f05.s1 Stratagene lung (#937210) H saplens cDNA clone IMAGE:	
				3' similar to contains Alu repetitive element; contains MER12 repetitive element;	
				mRNA sequence.	0.069
	100756	HG3565-HT3768		Zinc Finger Protein (Gb:M88357)	0.069
10	113483	T87768	Hs.16439	ESTs	0.069
10	101119	L09708	Hs.2253	complement component 2	0.069
	102286	U31628	Hs.12503	interleukin 15 receptor; alpha	0.07
			Hs.9930	collagen-binding protein 2 (colligen 2)	0.07
	135349	D83174			0.07
1.5	100991	J03764	Hs.82085	plasminogen activator inhibitor; type I	0.07
15	133675	AA443720	Hs.7551	ESTs; Weakly similar to T25G3.1 [C.elegans]	
	105422	AA251014	Hs.12210	ESTs	0.07
	102932	X13334	Hs.75627	CD14 antigen	0.07
	119147	R58878	Hs.65739	ESTs	0.07
	104900	AA055048	Hs.180481	ESTs; Weakly similar to ACROSIN PRECURSOR [H.sapiens]	0.07
20	133185	AA481404	Hs.6686	ESTs	0.07
-	115496	AA290674	Hs.71819	eukaryotic translation initiation factor 4E binding protein 1	0.07
	121005	AA398332	Hs.97613	ESTs	0.07
	124869	R69088	Hs.28728	ESTs; Weakly similar to F55A12.9 [C.elegans]	0.071
	129154	N23673	Hs.108969	mannosidase; alpha; class 2B; member 1	0.071
25	112161	R48295	110.110000	ESTs; Wkly smir to !! ALU SUBFAMILY J WARNING ENTRY !! [H.sapiens]	0.071
LJ	125251	W87486	Hs.141464	ESTs	0.071
•		J00116	Hs.81343	collagen; type II; alpha 1 (primary osteoarthritis; spondyloepiphyseal	
	134298	300110	113.01040		0.071
	440=45		11- 50000	dysplasia; congenital)	0.071
20	119745	W70264	Hs.58093	ESTs	0.071
30	131306	AA232686	Hs.25489	ESTs	
	107776	AA018820	Hs.221147	ESTs	0.071
	134271	AA199630	Hs.184456	ESTs; Wkty smlr to !! ALU SUBFAMILY SX WARNING ENTRY !! [H.sapiens]	0.071
	101798	M85220		Accession not listed in Genbank	0.071
	135402	S76942	Hs.99922	dopamine receptor D4	0.071
35	118742	N74052	Hs.50424	EST ·	0.071
	131867	N64656	Hs.3353	Homo sapiens clone 24940 mRNA sequence	0.071
	102923	X12517	Hs.1063	small nuclear ribonucleoprotein polypeptide C	0.072
	100775	HG371-HT26388		Mucin 1, Epithelial, Alt. Splice 9	0.072
	111020	N54361	Hs.185726	ESTs	0.072
40	134224	X80822	Hs.163593	ribosomal protein L18a	0.072
	124059	F13673	Hs.99769	ESTs	0.072
	133972	AA160743	Hs.78019	Homo sapiens clone 24432 mRNA sequence	0.072
	129681	AA436009	Hs.178186	ESTs; Weakly similar to WASP-family protein [H.sapiens]	0.072
		X58399	Hs.81221	Human L2-9 transcript of unrearranged immunoglobulin V(H)5 pseudogene	0.072
45	103065			calnexin	0.072
43	124966	T19271	Hs.155560	ESTs	0.072
	112270	R53021	Hs.203358		0.072
	116704	F10183	Hs.66140	EST	
	129890	M13699	Hs.111461	ceruloplasmin (ferroxidase)	0.072
	127345	AA972008	Hs.166253	ESTs; Highly similar to KIAA0476 protein [H.saplens]	0.072
50	112436	R63090	Hs.28391	ESTs	0.072
	114531	AA053033	Hs.203330	ESTs	0.072
	135122	H99080	Hs.94814	ESTs	0.072
	103934	AA281338	Hs.134200	Homo sapiens mRNA; cDNA DKFZp564C186 (from clone DKFZp564C186)	0.072
	109363	AA215369	Hs.185764	ESTs; Weakly similar to hypothetical protein [H.sapiens]	0.072
55	112647	R83329	Hs.33403	ESTs	0.073
	127083	Z44079	Hs.91608	otoferlin	0.073
	133027	AA402624	Hs.63236	synuclein; gamma (breast cancer-specific protein 1)	0.073
	122086	AA432121	Hs.250986	EST	0.073
		H47542	Hs.33962	ESTs	0.073
60	110405	AB002344	Hs.103915	KIAA0346 protein	0.073
JU	128697	R50380		ESTs	0.073
	112221		Hs.25670		0.073
	100478	HG1067-HT1067	LI- OFTOT	Mucin (Gb:M22406)	0.073
	115598	AA400129	Hs.65735	ESTs .	
	132491	AA227137	Hs.4984	KIAA0828 protein	0.073
65	101655	M60299		Human alpha-1 collagen type II gene, exons 1, 2 and 3	0.073
	106018	AA411887	Hs.34737	ESTs	0.073
	129683	W05348	Hs.158196	DKFZP434B103 protein	0.073
	134137	F10045	Hs.79347	KIAA0211 gene product	0.073
	114008	W89128	Hs.19872	ESTs	0.073
					P.
				199	•

	107653	AA010210	Hs.47041	ESTs	0.073
	104798	AA029462	Hs.17235	ESTs	0.073
	134082	L16991	Hs.79006	deoxythymidylate kinase	0.073
	119180	R80413	Hs.92520	ESTs	0.073
5	107741	AA016982	Hs.64341	ESTs	0.073
_	133683	AA335223	Hs.75558	pepsinogen 5; group I (pepsinogen A)	0.073
	111694	R22035	Hs.23331	ESTs	0.073
		AA338729	Hs.133096	ESTS	0.073
	120764				0.074
10	119389	T88826	Hs.90973	ESTs	0.074
10	100929	HG688-HT688		Major Histocompatibility Complex, Class II, Dr Beta 2 (Gb:X65561)	
	119388	T88798		plasminogen activator inhibitor, type I	0.074
	133019	AF009674	Hs.184434	axin	0.074
	105185	AA191495	Hs.189937	ESTs	0.074
	133413	S72043	Hs.73133	metallothionein 3 (growth inhibitory factor (neurotrophic))	0.074
15	101017	J04599	Hs.821	biglycan	0.074
	132865	K02765	Hs.251972	complement component 3	0.074
	110882	N36001	Hs.17348	ESTs; Wkly smir to !! ALU SUBFAMILY SQ WARNING ENTRY !! [H.sapiens]	0.074
	129197	T90303	Hs.109308	ESTs; Wkty smlr to leucine-rich glioma-inactivated prot precursor [H.sapiens]	0.074
	101184	L19871	Hs.460	activating transcription factor 3	0.075
20	134910	AA431320	Hs.9100	ESTs	0.075
20	119411	T96621	Hs.203656	EST	0.075
		U01824	Hs.380	solute carrier family 1 (glial high affinity glutamate transporter); member 2	0.075
	102000			ESTs; Weakly similar to envelope protein [H.sapiens]	0.075
	114691	AA121893	Hs.103779	plectin 1; intermediate filament binding protein; 500kD	0.075
25	134179	U53204	Hs.79706		0.075
25	134503	U34880	Hs.84183	diptheria toxin resistance protein required for diphthamide	0.076
			11 400000	biosynthesis (Saccharomyces)-like 1	0.075
	129719	N66396	Hs.167766	ESTs; Moderately similar to Pro-a2(XI) [H.sapiens]	0.075
	113916	W80464	Hs.31928	ESTs; Wkly smlr to alternatively spliced product using exon 13A [H.sapiens]	0.075
	113897	W73926	Hs.4947	ESTs	0.075
30	129697	R00841	Hs.172069	DKFZP434C212 protein	0.075
	112078	R44155	Hs.112218	ESTs	0.075
	121980	AA429886	Hs.110407	ESTs; Weakly similar to coded for by C. elegans cDNA yk173c12.5 [C.elegans]	0.075
	100898	HG4638-HT5050		Spliceosomal Protein Sap 49	0.075
	121626	AA416974	Hs.98174	ESTs	0.075
35	133670	AA243416	Hs.75470	hypothetical protein; expressed in osteoblast	0.075
	131879	AA017161	Hs.33792	ESTs	0.075
	100254	D38037	Hs.77643	FK506-binding protein 1B (12.6 kD)	0.075
	133194	AA291726	Hs.67201	ESTs	0.075
	106081	AA418394	Hs.25354	ESTs	0.075
40	115544	AA351433	Hs.66187	Homo sapiens clone 23700 mRNA sequence	0.076
40	119955	W87460	Hs.58989	ESTs ·	0.076
	104407	H61361	Hs.102171	immunoglobulin superfamily containing feucine-rich repeat	0.076
	135019	X58431	Hs.98428	Human Hox2.2 gene for a homeobox protein	0.076
			Hs.103931	DKFZP434B0335 protein	0.076
45	114815	AA161488 W31352	Hs.55445	ESTs	0.076
43	119471			ESTs	0.076
	117788	N48292	Hs.46849		0.076
	119406	T95064	Hs.193771	EST	0.076
	130777	R61742	Hs.256554	ESTs	
50	130494	L13197	Hs.75874	pregnancy-associated plasma protein A	0.076 0.076
50	104107	AA424111	Hs.12598	T-cell lymphoma invasion and metastasis 2	
	121483	AA411981	Hs.25274	ESTs; Modly smlr to putative seven pass transmembrane prot [H.sapiens]	0.076
	104451	M13299	Hs.102119	blue cone pigment	0.076
	118027	N52770	Hs.75968	thymosin; beta 4; X chromosome	0.076
	109419	AA227560	Hs.86987	receptor-interacting serine-threonine kinase 3	0.076
55	115783	AA424487	Hs.72289	ESTs; Weakly similar to LIV-1 protein [H.sapiens]	0.076
	110585	H62223	Hs.133526	ESTs; Wkly smir to !!!ALU SUBFAMILY SB1 WARNING ENTRY !!![H.sapiens]	0.076
	123165	AA488863	Hs.105216	ESTs; Weakly smir to !!ALU SUBFAMILY J WARNING ENTRY !! [H.sapiens]	0.077
	103966	AA303166	Hs.127270	ESTs	0.077
	109549	F01528	Hs.21192	Homo sapiens clone 25155 mRNA sequence	0.077
60	106730	AA465520	Hs.22313	ESTs	0.077
50	120310	AA193676	Hs.118926	DKFZP586K0919 protein	0.077
	104078	AA402801	Hs.222010	ESTs	0.077
		N35978	Hs.82364	ESTS	0.077
	117624			ESTS	0.077
65	112421	R62441	Hs.23127		0.077
65	106958	AA497026	Hs.22059	ESTS	
,	129984	W92811	Hs.183927	ESTs; Weakly similar to !! ALU SUBFAMILY J WARNING ENTRY !! [H.saplens]	0.077
	122044	AA431456	Hs.98736	EST	0.077
	123280	AA491285	Hs.175144	ESTs	0.077
	115710	AA412535	Hs.55235	sphingomyelin phosphodiesterase 2; neutra	

				harantara (a colon) ankia namo alianga)	0.077
		007111		I membrane (neutral sphingomyelinase)	0.077 0.077
	134129	D87444	Hs.79305	KIAA0255 gene product	
	129321	AA224502	Hs.206501	Homo sapiens clone 643 unknown mRNA; complete sequence	0.078
~	130513	AA460257	Hs.15866	ESTs	0.078
5	100996	J03909	Hs.14623	interferon; gamma-inducible protein 30	0.078
	128358	AI095718	Hs.135015	ESTs	0.078
	128544	R59352	Hs.119273	KIAA0296 gene product	0.078
	106040	AA412681	Hs.125139	ESTs	0.078
	106495	AA452113	Hs.32454	ESTs; Moderately similar to KIAA0544 protein [H.sapiens]	0.078
10	131833	R40899	Hs.32973	glycine receptor; beta	0.078
	119219	R97176	Hs.110783	ESTs	0.078
•	135415	X60655	Hs.99967	even-skipped homeo box 1 (homolog of Drosophila)	0.078
	109457	AA232646	Hs.68061	ESTs; Weakly similar to sphingosine kinase [M.musculus]	0.078
	117137	H96670	Hs.42221	ESTs ·	0.078
15	107094	AA609614	Hs.5241	ESTs	0.078
	130165	T90529	Hs.251613	EST	0.078
	124072	H05252	Hs.101637	EST; Weakly similar to hypothetical protein [H.sapiens]	0.078
	126151	AA324743	Hs.40808	ESTs	0.078
	119035	R01779	Hs.7740	ESTs	0.078
20	110157	H18987	Hs.169731	ESTs	0.078
	128515	AA149044	Hs.10086	ESTs; Highly similar to HYPOTHETICAL PROTEIN KIAA0195 [H.sapiens]	0.078
	133069	U94836	Hs.6430	protein with polyglutamine repeat	0.078
	112209	R49644	Hs.24865	ESTs	0.078
	133361	R28279	Hs.71848	Human clone 23548 mRNA sequence	0.078
25	134714	U89922	Hs.890	lymphotoxin beta (TNF superfamily; member 3)	0.078
	129905	T86796	Hs.132875	ESTs; Weakly similar to predicted using Genefinder [C.elegans]	0.079
	120421	AA236166	Hs.132957	ESTs; Weakly similar to chondromodulin-I precursor [H.sapiens]	0.079
	100885	HG4490-HT4876		Proline-Rich Protein Prb4, Allele	0.079
	102789	U86759	Hs.158336	netrin 2 (chicken)-like	0.079
30	120139	Z39273	Hs.77876	Human DNA from chromosome 19-specific cosmid R30923; genomic sequence	0.079
	135238	U76343	Hs.96970	Human liver GABA transport protein mRNA; 3' end	0.079
	129618	N54845	Hs.173030	ESTs	0.079
	132960	AA609742	Hs.6150	KIAA0521 protein	0.079
	108751	AA127063	Hs.203717	ESTs	0.079
35	134060	D42039	Hs.78871	KIAA0081 protein	0.079
	111338	N79778	Hs.35094	extracellular matrix protein 2; female organ and adipocyte specific	0.079
	112345	R56880	Hs.26563	ESTs	0.079
	126456	W00881		za56d02.r1 Soares fetal liver spleen 1NFLS Homo sapiens cDNA done	0.000
			•	IMAGE:296547 5', mRNA sequence.	0.079
40	128937	Z39939	Hs.10726	ESTs	0.079
	103485	Y08409 ·	Hs.248415	thyroid hormone responsive SPOT14 (rat) homolog	0.079
	111202	N68280	Hs.107922	ESTs	0.079
	132625	AA429890	Hs.166066	cisplatin resistance associated	0.079
	103434	X98085	Hs.54433	tenascin R (restrictin; janusin)	0.079
45	102616	U65581	Hs.159191	ribosomal protein L3-like	0.079
	102667	U70867	Hs.83974	solute carrier family 21 (prostaglandin transporter); member 2	0.079
	111422	R01127	Hs.19104	ESTs	0.079
	101411	M16938	Hs.820	homeo box C6	80.0
~ 0	113267	T65058	Hs.12725	ESTs; Weakly similar to !! ALU SUBFAMILY J WARNING ENTRY !! [H.sapiens]	80.0
50	103559	Z19585	Hs.75774	thrombospondin 4	80.0 80.0
	131588	AA258613	Hs.29189	KIAA1021 protein	0.08
	107821	AA020991	Hs.172856	ESTs	0.08
	134278	H82839	Hs.81001	ESTs; Weakly similar to DY3.6 [C.elegans]	
~ ~	120893	AA369800	Hs.97058	EST; Highly similar to CMP-N-acetylneuraminic acid hydroxylase [H.sapiens]	80.0
55	108786	AA128999		zo8f12.s1 Stratagene neuroepithelium NT2RAMI 937234 Homo sapiens	80.0
				cDNA clone IMAGE:567119 3', mRNA sequence	0.08
	106890	AA489245	Hs.88500	KIAA1066 protein; JSAP1 homolog (mouse); JIP3 homolog (mouse)	0.08
	119760	W72267	Hs.58219	ESTs	0.08
60	132999	Y00787	Hs.624	interleukin 8	0.08
60	129156	AA028195	Hs.108973	dolichyl-phosphate mannosytransferase polypeptide 2; regulatory subunit	0.08
	121171	AA400008	Hs.161814	ESTs	0.08
	103864	AA207264	Hs.181077	ESTs; Weakly similar to Miller-Dieker lissencephaly gene [H.sapiens]	0.08
	128591	AA255537	Hs.102057	ESTs; Weakly similar to O-linked GlcNAc transferase [H.sapiens]	0.08
15	122172	AA435753	Hs.161854	EST	0.08
65	112802	R97647	Hs.174855	EST	0.08
	107723	AA015967	Hs.60680	EST	0.081
	113011	T23737	Hs.1600	chaperonin containing TCP1; subunit 5 (epsilon)	0.081
	131279	AA089853	Hs.25197	STIP1 homology and U-Box containing protein 1	0.081
	103190	X70083	Hs.58414	filamin C; gamma (actin-binding protein-280)	U.UU I

				•	
	103956	AA292411	Hs.233348	ESTs	0.081
	112706	R89828	Hs.138493	ESTs	0.081
	126126	M85370		EST01884 Fetat brain, Stratagene (cat#936206) Homo sapiens cDNA	
		•		clone HFBCH10, mRNA sequence.	0.081
5	130094	H43286	Hs.167017	gamma-aminobutyric acid (GABA) B receptor; 1	0.081
_	100800	HG3945-HT4215		Phospholipid Transfer Protein	0.081
	108675	AA115240	Hs.61816	ESTs	0.081
	129420	AA234259	Hs.99816	ESTs	0.081
	129666	M77349	Hs.118787	transforming growth factor; beta-induced; 68kD	0.081
10	101645	M59807	Hs.943	natural killer cell transcript 4	0.081
	130536	T17045	Hs.159492	spastic ataxia of Charlevoix-Saguenay (sacsin)	0.081
	107732	AA016181	Hs.59752	ESTs	0.081
	123071	AA482593	Hs.104285	ESTs .	0.081
	113537	T90457	Hs.191293	ESTs	0.081
15	101250	L34060	Hs.79133	cadherin 8	0.081
	122521	AA449433	Hs.149227	ESTs; Weakly similar to PROLINE-RICH PROTEIN MP-3 [M.musculus]	0.081
	133914	N32811	Hs.77542	ESTs	0.081
	102038	U05659	Hs.477	hydroxysteroid (17-beta) dehydrogenase 3	0.081
00	110336	H40338	Hs.174094	ESTs; Weakly similar to !! ALU SUBFAMILY J WARNING ENTRY !! [H.sapiens]	0.081
20	118637	N70274	Hs.49822	ESTs	0.081
	117966	N51589	Hs.94012	ESTs	0.082 0.082
	104424	H87671	Hs.182320	ESTs; Weakly similar to Mouse 19.5 mRNA; complete cds [M.musculus]	0.082
	100361	D78361	Hs.125078	Human mRNA for omithine decarboxylase antizyme; ORF 1 and ORF 2 microtubule-associated protein tau	0.082
25	112974	T17291 D63482	Hs.101174 Hs.57734	KIAA0148 gene product	0.082
23	132832 132039	Z39489	Hs.3781	Homo sapiens BAC clone RG118D07 from 7q31	0.082
	113272	T65383	Hs.12807	ESTs	0.082
	104924	AA058532	Hs.28774	ESTs	0.082
	111061	N58054	Hs.36859	ESTs	0.082
30	129269	R45977	Hs.163593	ribosomal protein L18a	0.082
	102453	U48437	Hs.74565	amyloid beta (A4) precursor-like protein 1	0.082
	126204	AI080388	Hs.134296	ESTs	0.082
	116615	D80666	Hs.45203	ESTs	0.082
	128856	AA219552	Hs.204144	ESTs; Modly smir to tumor necrosis factor-alpha-induced prot B12 [H.sapiens]	0.082
35	112776	R95850	Hs.34494	ESTs	0.082
	105494	AA256273	Hs.29288	Homo sapiens mRNA; cDNA DKFZp434P174 (from clone DKFZp434P174)	0.082
	117000	H84718	Hs.112236	ESTs; Weakly similar to repressor protein [H.sapiens]	0.082 0.082
	112656	R85260	Hs.133151	transient receptor potential channel 7	0.083
40	128963	J03890 H79292	Hs.1074 Hs.39960	surfactant; pulmonary-associated protein C ESTs	0.083
40	116957 101057	K03430	115.03300	Human complement C1g B-chain gene, exon A+1	0.083
	121948	AA429452	Hs.98582	ESTs	0.083
	130822	M80647	Hs.2001	thromboxane A synthase 1 (platelet; cytochrome P450; subfamily V)	0.083
	122743	AA458674	Hs.99478	EST	0.083
45	114569	AA063316	*	zm2d1.s1 Stratagene corneal stroma (#937222) Homo sapiens cDNA clone	
				IMAGE:512947 3' similar to TR:E198281 E198281 THIOREDOXIN	
				REDUCTASE ;contains Alu repetitive element;, mRNA sequence	0.083
	132270	U70671	Hs.43509	ataxin 2 related protein	0.083
	108126	AA052951	Hs.47413	ESTs	0.083
50	102880	X04325	Hs.2679	gap junction protein; beta 1; 32kD (connexin 32; Charcot-Marie-Tooth	0.000
			11 00500	neuropathy; X-linked)	0.083
	115365	AA282089	Hs.88599	ESTS	0.083
	114529	AA052980	Hs.206704	ESTs - ESTs; Weakly similar to NEURONAL OLFACTOMEDIN-RELATED	0.003
55	135017	AA249586	Hs.9315	ER LOCALIZED PROTEIN [H.sapiens]	0.083
33	123776	AA610071	Hs.112813	ESTs	0.083
	114454	AA021091	Hs.226208	ESTs	0.083
	101246	L33799	Hs.202097	procollagen C-endopeptidase enhancer	0.083
	107366	U78310	Hs.13501	pescadillo (zebrafish) homolog 1; containing BRCT domain	0.083
60	132779	T89601	Hs.95497	ESTs; Weakly similar to GLUCOSE TRANSPORTER TYPE 5;	
-				SMALL INTESTINE [H.sapiens]	0.083
	129709	AA112209	Hs.1209	acyl-Coenzyme A dehydrogenase; long chain	0.083
	115244	AA278767	Hs.914	Human mRNA for SB classil histocompatibility antigen alpha-chain	0.083
<i></i>	123253	AA490878	Hs.111334	ferritin; light polypeptide	0.083
65	128469	T23724	Hs.258677	EST	. 0.083
	132220	AA431847	Hs.42409	ESTs; Highly similar to CGI-146 protein [H.saplens]	0.083
	111664	R17939	Hs.22344	ESTs	0.083
	102354	U38268	11-404000	Human cytochrome b pseudogene, partial cds	0.084 0.084
	112828	R98774	Hs.194338	ESTs	V.U04

	110410	H47868	Hs.34024	ESTs	0.084
	102620	U66052		Human clone W2-6 mRNA from chromosome X	0.084
	102550	U58087	Hs.14541	cullin 1	0.084
	108417	AA075716		zm89e5.s1 Stratagene ovarian cancer (#937219) H sapiens cDNA clone	
5				IMAGE:54512 3' similar to gb:X14723 CLUSTERIN PRECURSOR	
_				(HUMAN);, mRNA sequence	0.084
•	113299	T67285	Hs.13089	ESTs	0.084
	117869	N49947	Hs.46990	ESTs	0.084
	113734	T98484	Hs.18377	EST	0.084
10	133325	C00424	Hs.7101	periodontal ligament fibroblast protein	0.084
	123368	AA505022	Hs.124838	ESTs	0.084
	101615	M55153	Hs.8265	transglutaminase 2 (C polypeptide; protein-glutamine	
				-gamma-glutamyttransferase)	0.084
	119352	T65972	Hs.193365	ESTs; Moderately similar to alternatively spliced product	
15				using exon 13A [H.sapiens]	0.084
	123828	AA620686	Hs.112884	EST	0.084
	103611	Z38133	Hs.113973	myosin; heavy polypeptide 8; skeletal muscle; perinatal	0.084
	131289	AA485697	Hs.25334	ESTs; Weakly similar to ION CHANNEL HOMOLOG RIC	
				PRECURSOR [M.musculus]	0.084
20	128678	T15896	Hs.103535	ESTs	0.084
	130814	AA256695	Hs.19813	ESTs	0.084
	133391	X57579	Hs.727	inhibin; beta A (activin A; activin AB alpha polypeptide)	0.084
	129322	AA437153	Hs.110407	ESTs; Weakly similar to coded for by C. elegans cDNA yk173c12.5 [C.elegans]	0.084
	109284	AA196995	Hs.86092	ESTs	0.084
25	116689	F09222	Hs.66099	ESTs	0.085
	100545	HG2147-HT2217		Mucin 3, Intestinal (Gb:M55405)	0.085
	102634	U66711	Hs.77667	lymphocyte antigen 6 complex; locus E	0.085
	111735	R25389	Hs.23856	ESTs; Weakly similar to FAST kinase [H.saplens]	0.085
	105181	AA190676	Hs.10974	ESTs; Moderately similar to unknown [R.norvegicus]	0.085
30	122681	AA455350	Hs.99401	EST	0.085
	114543	AA056121	Hs.158419	ESTs	0.085
	133597	AA425908	Hs.75139	partner of RAC1 (arfaptin 2)	0.085
	121064	AA398647	Hs.97406	ESTs	0.085
0.5	122231	AA436369	Hs.197728	ESTs; Weakly similar to ZINC FINGER PROTEIN 135 [H.sapiens]	0.085
35	100309	D50550	Hs.95659	lethal giant larvae (Drosophila) homolog 1	0.085
	101727	M73481	Hs.73883	gastrin-releasing peptide receptor	0.085
	131226	AA165400	Hs.24476	ESTs	0.085
	133580	AA095041	Hs.181073	ESTs	0.085 0.085
40	102792	U87964	Hs.227576	GTP binding protein 1	0.085
40	104976	AA086480	Hs.183669	ESTs; Weakly similar to !! ALU SUBFAMILY J WARNING ENTRY !! [H.sapiens]	0.085
	120865	AA350631	Hs.96963	EST	0.085
	106080	AA418046	Hs.35124	ESTs	0.085
	128571	AA416619	Hs.101661	ESTs connective tissue growth factor	0.085
45	101838	M92934	Hs.75511 Hs.100843	ESTs; Weakly similar to similar to GTP-binding protein [C.elegans]	0.085
43	128514 123099	H84261 AA485931	Hs.79	aminoacylase 1	0.085
	134067	Y08200	Hs.78920	Rab geranyigeranyitransferase; alpha subunit	0.085
	116967	H80336	Hs.40124	EST	0.085
	110053	H12586	Hs.89563	nuclear cap binding protein 1; 80kD	0.085
50	114395	AA007313	Hs.110155	ESTs	0.085
50	107465	W44681	Hs.251385	murine retrovirus Integration site 1 homolog	0.085
	101983	S85655	Hs.75323	prohibitin	0.085
	112544	R70948	Hs.29153	ESTs -	0.086
	111423	R01165	Hs.188507	ESTs .	0.086
55	127918	AA806043	Hs.115396	Human germline IgD chain gene; C-region; C-delta-1 domain	0.086
23	107300	T40348	Hs.90488	ESTs	0.086
	134947	R51194		vi71a08.r1 Soares breast 2NbHBst Homo sapiens cDNA clone IMAGE:154166	
				5' similar to gb:L11284 DUAL SPECIFICITY MITOGEN-ACTIVATED PROTEIN	
		•		KINASE KINASE 1 (HUMAN);, mRNA sequence.	0.086
60	124579	N68345	Hs.127179	ESTs; Weakly similar to TERATOCARCINOMA-DERIVED GROWTH	
			•	FACTOR 1 [H.sapiens]	0.086
	130471	Z68280	Hs.183706	adducin 1 (alpha)	0.086
	116596	D60755	Hs.92955	ESTs	0.086
	105069	AA136345	Hs.23617	ESTs; Weakly similar to ZFOC1 gene product [H.sapiens]	0.086
65	102491	U51010		Human nicotinamide N-methyltransferase gene, exon 1 and 5' flanking region	0.086
	130069	AA055896	Hs.146428	collagen; type V; alpha 1	0.086
	130234	AA280413	Hs.157441	spleen focus forming virus (SFFV) proviral integration oncogene spi1	0.086
	120540	AA262992	Hs.96417	ESTs	0.086
	122508	AA449221	Hs.20432	ESTs	0.086

	128054	Al205718	Hs.125416	ESTs	0.086
	133020	AA053248	Hs.185182	ESTs; Highly similar to 40S RIBOSOMAL PROTEIN S10 [H.sapiens]	0.086
	130056	AA017356	Hs.171900	armadillo repeat gene deletes in velocardiofacial syndrome	0.086
	130504	U48865	Hs.158323	CCAAT/enhancer binding protein (C/EBP); epsilon	0.086
5	133978	W73859	Hs.78061	transcription factor 21	0.086
	105265	AA227941	Hs.26088	ESTs	0.086
	133035	T15965	Hs.6333	ESTs	0.086
	100768	HG3636-HT3846		Myosin, Heavy Polypeptide 9, Non-Muscle	0.086
	129338	T56800	Hs.47274	Homo sapiens mRNA; cDNA DKFZp564B176 (from clone DKFZp564B176)	0.086
10	132789	W23761	Hs.56876	ESTs	0.086
	116099	AA456309	Hs.58831	regulator of Fas-Induced apoptosis	0.086
	100721	HG3355-HT3532		Peroxisome Proliferator Activated Receptor (Gb:Z30972)	0.087
	112569	R73150	Hs.75270	GTP-binding protein homologous to Saccharomyces cerevisiae SEC4	0.087
	130645	AA020942	Hs.17200	STAM-like protein containing SH3 and ITAM domains 2	0.087
15	100751	HG3527-HT3721		Luteinizing Hormone, Beta Subunit	0.087
	134550	M27161	Hs.85258	CD8 antigen; alpha polypeptide (p32)	0.087
	130885	AA338646	Hs.20912	adenomatous polyposis coli like	0.087
	101446	M21302	Hs.56306	small proline-rich protein 2A	0.087
	116287	AA487856	Hs.155829	KIAA0676 protein	0.087
20	134034	X89267	Hs.78601	uroporphyrinogen decarboxylase	0.087
	130860	U66061	Hs.241395	protease; serine; 1 (trypsin 1)	0.087
	109901	H04992	Hs.30499	ESTs	0.087
	107537	Z20777	Hs.9857	ESTs; Weakly similar to peroxisomal short-chair alcohol	
				dehydrogenase [H.sapiens]	0.087
25	133232	AA496030	Hs.6845	ESTs	0.087
	108559	AA085161		zn12c5.s1 Stratagene hNT neuron (#937233) H sapiens cDNA clone	
				IMAGE:54728 3' similar to TR:G1151228 G1151228 LPG1P.;, mRNA seq	0.087
	121288	AA401735	Hs.97340	EST	0.087
	108844	AA132916	Hs.177961	Human Chromosome 16 BAC done CIT987SK-A-388D4	0.087
30	129874	AA406488	Hs.181551	ESTs	0.087
	105139	AA164543	Hs.110082	ESTs	0.088
	124789	R43803	Hs.78110	ESTs; Weakly similar to F17A9.2 [C.elegans]	0.088
	115923	AA441929	Hs.38205	ESTs	0.088
25	123640	AA609292	Hs.112681	ESTs	0.088
35	131607	AA351409	Hs.172740	microtubule-associated protein; RP/EB family; member 3	0.088
	130064	T67053	Hs.181125	immunoglobulin lambda gene cluster	880.0
	108752	AA127070	Hs.71055	ESTs	0.088
	124249	H68077	Hs.108211	ESTs	0.088
40	100109	AJ000480	Hs.143513	phosphoprotein regulated by mitogenic pathways	0.088
40	104642	AA004662	Hs.184245	KIAA0929 protein Msx2 interacting nuclear target (MINT) homolog	0.088 0.088
	131752	AA453311	Hs.31566	ESTs	0.088
	114727 120965	AA132545 AA398089	Hs.190202 Hs.179715	ESTs ESTs	0.088
	100396	D84361	Hs.151123	Human mRNA for p52 and p64 isoforms of N-Shc; complete cds	0.088
45	106218	AA428451	Hs.91146	DKFZP586E0820 protein	0.088
43	111562	R09567	Hs.187569	ESTs	0.088
	121219	AA400606	Hs.144344	EST	0.088
	101187	L20316	Hs.208	glucagon receptor	0.088
	101513	M28210	Hs.27744	RAB3A; member RAS oncogene family	0.088
50	116454	AA621071	Hs.42034	ESTs; Moderately similar to T-complex protein 10A [H.sapiens]	0.088
50	116171	AA463434	Hs.42658	ESTs	0.089
	117500	N31909	Hs.44278	ESTs	0.089
	119978	W88623	Hs.59190	EST -	0.089
	132005	D58231	Hs.173091	DKFZP434K151 protein	0.089
55	109914	H05529	Hs.194704	leucine-rich; glioma inactivated 1	0.089
	130370	M55265	Hs.155140	casein kinase 2; alpha 1 polypeptide	0.089
	104262	AF009801	Hs.105941	bagpipe homeobox (Drosophila) homolog 1	0.089
	129708	AA417181	Hs.120858	ESTs	0.089
	106398	AA447545	Hs.18268	adenylate kinase 5	0.089
60	120884	AA365356	Hs.97041	ESTs	0.089
-	130404	X72012	Hs.76753	endoglin (Osler-Rendu-Weber syndrome 1)	0.089
	114072	Z38184	Hs.123633	ESTs	0.089
	131470	X54938	Hs.2722	inositol 1;4;5-trisphosphate 3-kinase A	0.089
	124573	N67935	Hs.194703	adaptor-related protein complex 4; mu 1 subunit	0.089
65	114717	AA131240	Hs.252014	EST	0.089
	133806	M12759	Hs.76325	Human 1g J chain gene	0.09
	130470	AA398552	Hs.15711	KIAA0639 protein	0.09
	133182	Z80787	Hs.240135	H4 histone family; member J	0.09
	116036	AA452572	Hs.43866	ESTs .	0.09

	132404	AA393903	Hs.4768	ESTs	0.09
	122695	AA456048	Hs.99403	ESTs; Moderately similar to undulin 2 [H.sapiens]	0.09
	125975	AA495891	Hs.152290	ESTs; Highly similar to PACAP type-3/VIP type-2 receptor [H.sapiens]	0.09
_	110783	N23669	Hs.26407	ESTs	0.09
5	129860	AA410343	Hs.129826	tetraspan transmembrane 4 super family	0.09
	120740	AA302650	Hs.96654	EST	0.09
	119564	W38206		Accession not listed in Genbank	0.09
	134474	AA054746	Hs.8379	ESTs	0.09
	119014	N95435	Hs.55144	ESTs	0.09
10	109791	F10669	Hs.13228	DRE-antagonist modulator; calsenilin	0.09
	117605	N35073	Hs.44433	ESTs	0.09
	121589	AA416627	Hs.191598	ESTs	0.09
	104326	D81655	Hs.143067	ESTs	0.09
	129861	N69507	Hs.129849	DKFZP564M182 protein	0.09
15	102795	U88667	Hs.198396	ATP-binding cassette; sub-family A (ABC1); member 4	0.09
	119626	W49499	Hs.184456	ESTs; Wkty smir to II ALU SUBFAMILY SX WARNING ENTRY !! [H.sapiens]	0.09
	110516	H56894	Hs.37368	EST	0.09
	105382	AA236853	Hs.111801	Homo sapiens mRNA; cDNA DKFZp564H2023 (from clone DKFZp564H2023)	0.09
	123754	AA609964	Hs.102021	ESTs	0.09
20	108008	AA039430	Hs.61920	ESTs	0.09
20	121057	AA398619	Hs.142375	ESTs; Moderately similar to putative envelope protein [H.sapiens]	0.091
	123675	AA609474	Hs.112713	EST	0.091
	135194	C20975	Hs.9613	ESTs; Highly similar to angiopoletin-related protein [H.sapiens]	0.091
	127070	AA641812	Hs.190037	ESTs	0.031
25	134051	S67070	Hs.78846	heat shock 27kD protein 2	0.091
ديد	133382	AA112532	Hs.7247	ESTs	0.091
	103615	Z46967	Hs.115460	calicin	0.091
	118457	N66593	Hs.49230	EST	0.091
	118504	N67334	Hs.50158	ESTs	0.091
30	112915	T10176	Hs.4254	ESTs	0.091
50	132088	AA470121	Hs.243960	HLA-B associated transcript-3	0.091
	101504	M27288	Hs.248156	oncostatin M	0.091
	112550	R71391	Hs.29074	ESTs	0.091
	128551	H09058	Hs.237323	N-acetylglucosamine-phosphate mutase; DKFZP434B187 protein	0.091
35	112879	T03541	Hs.115960	ESTs	0.091
33	127079	Al364691	Hs.128628	ESTs; Moderately similar to CL3BC [R.norvegicus]	0.091
	101993	U01062	Hs.77515	inositol 1;4;5-triphosphate receptor; type 3	0.091
	113020	T23830	Hs.7303	ESTs; Weakly similar to PROHIBITIN [H.saplens]	0.091
	120465	AA251505	Hs.130861	ESTs. Weakly similar to Phohibit in [h.sapiens]	0.091
40	130152	U32645	Hs.151139	E74-like factor 4 (ets domain transcription factor)	0.091
40	104941	AA065169	Hs.17805	ESTs	0.091
	110090	H16076	Hs.6915	ESTs	0.091
	135375	AA480888	Hs.99741	ESTS; Weakly similar to BRAIN PROTEIN H5 [H.sapiens]	0.091
	123799	AA620418	Hs.112861	ESTs	0.092
45	118966	N93438	Hs.76907	ESTs; Highly similar to HSPC002 [H.sapiens]	0.092
73	116969	H80633	Hs.143038	ESTs	0.092
	125147	W38150	FIS. 140000	Accession not listed in Genbank	0.092
	100836	HG4113-HT4383		Olfactory Receptor 0r17-201	0.092
	114726	AA132509	Hs.103827	EST	0.092
50	107311	T57738	Hs.174112	ESTs	0.092
50			Hs.4610	EST	0.092
	112863 129290	T03148		ESTs	0.092
	103384	AA521407	Hs.110095	tafazzin (cardiomyopathy; dilated 3A (X-linked); endocardial	0.032
	100004	X92762	Hs.79021	fibroelastosis 2: Barth syndrome)	0.092
55	112508	R68213	Hs.28847	ESTs	0.092
33	111863	R37495	Hs.23578	ESTS	0.092
	131184	AA452705	Hs.23954	ESTs; Weakly similar to KIAA0584 protein [H.sapiens]	0.092
	107420	W26567	Hs.4775	ESTs, Weakly summar to KiAACOO4 protein [m.sapiens]	0.092
	111768	R27606		ESTs	0.092
60	112290		Hs.24185	ESTS ESTS	0.092
UU	130581	R53940	Hs.26016	ESTS; Weakly similar to RAS-RELATED PROTEIN RAB-5A [H.sapiens]	0.092
		AA481982	Hs.16258		
	120744	AA302772	Hs.228649	EST	0.093
	112226	R50761	Hs.25738	ESTs	0.093
65	116154	AA460951	Hs.57100	ESTs	0.093
O)	102640	U67674	Hs.194783	solute carrier family 10 (sodium/bile acid cotransporter family); member 2	0.093
	129797	X53595	Hs.1252	apolipoprotein H (beta-2-glycoprotein I)	0.093
	102705	U77180	Hs.50002	small Inducible cytokine subfamily A (Cys-Cys); member 19	0.093
	132408	AA035547	Hs.47822	KIAA0380 gene product; RhoA-specific guanine nucleotide exchange factor	0.093
	108441	AA079079		zm97c9.s1 Stratagene colon HT29 (#937221) Homo sapiens cDNA clone	

				IMAGE:545872 3' similar to contains element MER22 MER22 repetitive	
				element; mRNA sequence	0.093
	108145	AA054133	Hs.63085	ESTs	0.093
	106466	AA449990	Hs.76057	lysophospholipase II	0.093
5	101697	M64358		Human rhom-3 gene, exon	0.093
J	121294	AA401958	Hs.240170	ESTs; Moderately similar to alternatively spliced product using	
	121204	AA401330	113.240170	exon 13A [H.sapiens]	0.093
	117004	N49065	Hs.125201	ESTs; Weakly similar to B7 [M.musculus]	0.093
	117824	AA422049	Hs.40780	ESTs	0.093
10	115771	U33053	Hs.2499	protein kinase C-like 1	0.093
10	102303	U79255	Hs.26468	amyloid beta (A4) precursor protein-binding; family A; member 2 (X11-like)	0.093
	131405		Hs.101094	ESTs	0.093
	112909	T10069		ESTs	0.093
•	124173	H41281	Hs.107619	ESTS	0.093
15	112488	R66896	Hs.28788		0.093
13	130554	X59303	Hs.159637	valyHRNA synthetase 2	0.093
	106413	AA447964	Hs.6311	ESTs	0.093
	111711	R22891	Hs.7093	ESTs	0.094
	117595	N34933	Hs.44664	EST	0.094
20	113813	W45174	Hs.31382	ESTs	0.094
20	107769	AA018449	Hs.125220	Homo sapiens DNA from chromosome 19-cosmids R30102:R29350:R27740	0.004
				containing MEF2B; genomic sequence	0.094
	114966	AA250743	Hs.92198	ESTs; Highly similar to calcium-regulated heat stable protein	0.004
				CRHSP-24 [H.sapiens]	0.094
0.5	130297	H94949	Hs.171955	trophinin-assisting protein (tastin)	0.094
25	109589	F02429	Hs.6581	ESTs	0.094
	112592	R77631	Hs.29126	ESTs	0.094
	102314	U34038	Hs.154299	coagulation factor II (thrombin) receptor-like 1	0.094
	116128	AA459915	Hs.112193	mutS (E. coli) homolog 5	0.094
	106809	. AA479704	Hs.220324	Human DNA sequence from clone 283E3 on chromosome 1p36.21-36.33.	
30				Contains the alternatively spliced gene for Matrix Metalloproteinase in the	
				Female Reproductive tract MIFR1; -2; MMP21/22A; -B and -C; a novel gene;	0.004
				the alternatively spliced CDC2L2 gene for	0.094
	130607	AA043894	Hs.16603	ESTs	0.094
~-	120592	AA281929	Hs.143974	ESTs	0.094
35	117230	N20535	Hs.43265	melastatin 1	0.094
	105948	AA404597	Hs.7133	ESTs	0.094
	101333	L47738	Hs.80313	p53 inducible protein	0.094
	101909	S69265 .		Homo sapiens mRNA for PLE21 protein; complete cds	0.094
	106959	AA497031	Hs.8657	ESTs; Highly similar to CTG7a [H.sapiens]	0.094
40	127034	AA352389		ESTs; Wkly smlr to glucose-6-phosphatase catalytic subunit [R.norvegicus]	0.095
	134430	H52105	Hs.8309	KIAA0747 protein	0.095
	120342	AA207105	Hs.45068	Homo sapiens mRNA; cDNA DKFZp434l143 (from clone DKFZp434l143)	0.095
	104450	L77564	Hs.103978	serine/threonine kinase 22B (spermiogenesis associated)	0.095
4	130902	AA424530	Hs.21061	ESTs	0.095
45	102708	U77594	Hs.37682	retinoic acid receptor responder (tazarotene induced) 2	0.095
	107373	U85773	Hs.154695	phosphomannomutase 2	0.095
	123569	AA608952	Hs.195292	ESTs; Weakly similar to RNA helicase HDB/DICE1 [H.saplens]	0.095
	102687	U73379	Hs.93002	ubiquitin carrier protein E2-C	0.095
~~	128888	AA034951	Hs.106893	ESTs	0.095
50	100283	D43642	Hs.2430	transcription factor-like 1	0.095
	102747	U79303	Hs.82482	protein predicted by clone 23882	0.095
	107798	AA019346	Hs.60918	EST	0.095
	123565	AA608907	Hs.112614	EST -	0.095
	116010	AA449450	Hs.56421	ESTs; Weakly similar to Similarity to H.influenza ribonuclease PH [C.elegans]	0.095
55	117155	H97536	Hs.42391	EST	0.095
	133094	AA115572	Hs.64746	chloride Intracellular channel 3	0.095
	113174	T54659	Hs.9779	ESTs	0.095
	102016	U03270	Hs.122511	centrin; EF-hand protein; 1	0.095
	130126	AB002318	Hs.150443	KIAA0320 protein	0.095
60	134813	X14767	Hs.89768	gamma-aminobutyric acid (GABA) A receptor, beta 1	0.095
	132055	N69440	Hs.38132	ESTs	0.095
	122229	AA436198	Hs.103902	ESTs	0.096
	127574	AA907314	Hs.188905	ESTs .	0.096
	134432	AA053022	Hs.8312	ESTs	0.096
65	128052	AA878398	Hs.190491	ESTs	0.096
	101637	M58285	Hs.132834	hematopoletic protein 1	0.096
	103386	X92972	Hs.80324	protein phosphatase 6; catalytic subunit	0.096
	133079	AA477561	Hs.6449	ESTs	0.096
	120328	AA196979	Hs.104129	ESTs; Weakly similar to protease [H.sapiens]	0.096

	107640	AA009615	Hs.257808	ESTs	0.096
	123389	AA521176	Hs.221231	ESTs	0.096
	103222	· X74795	Hs.77171	minichromosome maintenance deficient (S. cerevisiae) 5 (cell division cycle 46)	0.096
	111704	R22450	Hs.23396	ESTs; Highly similar to ZINC FINGER PROTEIN 140 [H.saplens]	0.096
5	126856	AA306523		EST177475 Jurkat T-cells VI Homo sapiens cDNA 5' end, mRNA sequence.	0.733
-	127071	AA250806		ESTs	0.096
	114550	AA056755	Hs.151714	ESTs	0.096
	125955	Al356943	Hs.143761	ESTs	0.096
	134363	M37033	Hs.82212	CD53 antigen	0.096
10	128550	W76492	Hs.170142	ESTs	0.096
10	122598	AA453465	Hs.99329	ESTs	0.096
	118898	N90703	Hs.4236	KIAA0478 gene product	0.096
					0.096
	117661	N39092	Hs.44940	ESTs	0.096
15	120996	AA398281	Hs.143684	ESTs	0.096
13	123388	AA521172	Hs.134417	ESTs	0.096
	106700	AA463929	Hs.28701	ESTs	
	112962	T16814	Hs.6828	ESTS	0.096
	121262	AA401372	Hs.97723	ESTs	0.096
20	134551	R44839	Hs.8526	i-beta-1;3-N-acetylglucosaminyltransferase	0.096
20	112060	R43754	Hs.21164	ESTs	0.096
	134678	AA039935	Hs.182595	dynein; axonemal; light polypeptide 4	0.096
	100855	HG4234-HT4504		Methylenetetrahydrofolate Reductase	0.097
	132414	N91193	Hs.48145	ESIS	0.097
	112900	T08758	Hs.3813	ESTs	0.097
25	115989	AA447777	Hs.93135	ESTs	0.097
	103561	Z21488	Hs.143434	contactin 1	0.097
	131087	AA009738	Hs.22824	ESTs; Weakly similar to p160 myb-binding protein [M.musculus]	0.097
	120293	AA190859	Hs.191428	ESTs	0.097
	111830	R36081	Hs.25085	EST	0.097
30	113654	T95770	Hs.17666	ESTs	0.097
	132675	AA179338	Hs.5476	serine proteinase inhibitor	0.097
	120182	Z40125	Hs.91968	ESTs	0.097
	132879	U16282	Hs.5881	ELL gene (11-19 lysine-rich leukemia gene)	0.097
	134211	AA056681	Hs.80021	ESTs; Weakly similar to 62D9.p [D.melanogaster]	0.097
35	115448	AA284845	Hs.165051	ESTs	0.097
	118118	N56901	Hs.47995	ESTs	0.097
	107598	AA004528	Hs.169444	ESTs	0.097
	128933	H01824	Hs.760	GATA-binding protein 2	0.097
	114892	AA235988	Hs.86024	ESTs	0.097
40	101922	S75168	Hs.274	megakaryocyte-associated tyrosine kinase	0.097
	105444	AA252374	Hs.19333	ESTs; Weakly similar to ATP(GTP)-binding protein [H.sapiens]	0.097
	128155	AA926843	Hs.143302	ESTs	0.097
	116276	AA485870	Hs.44914	ESTs ·	0.097
	111964	R41227	Hs.21860	ESTs	0.097
45	135100	AA398926	Hs.251108	Homo sapiens mRNA; chromosome 1 specific transcript KIAA0493	0.097
	124872	R69251	Hs.101506	EST	0.097
	103084	X59932	Hs.77793	c-src tyrosine kinase	0.097
	124138	H23199	Hs.107010	ESTs	0.098
	130048	R31745	Hs.211612	SEC24 (S. cerevisiae) related gene family; member A	0.098
50	100208	D26129	Hs.78224	ribonuclease; RNase A family; 1 (pancreatic)	0.098
	123537	AA608775	Hs.112589	ESTs	0.098
	118999	N95019	Hs.55092	ESTs	0.098
	119847	W80384	Hs.9853	ESTs -	0.098
	112819	R98618	Hs.35984	ESTs	0.098
55	131080	J05008	Hs.2271	endothelin 1	0.098
	127353	AA190853	Hs.155360	ESTs	0.098
	132068	X66365	Hs.38481	cyclin-dependent kinase 6	0.098
	105744	AA293436	Hs.12909	ESTs	0.098
	133680	M92357	Hs.101382	tumor necrosis factor, alpha-induced protein 2	0.098
60	122899	AA469960	Hs.178420	ESTs; Highly similar to WASP Interacting protein [H.sapiens]	0.098
	128700	U59286	Hs.103982	small inducible cytokine subfamily B (Cys-X-Cys); member 11	0.098
	104393	H46486	Hs.226499	nesca protein	0.098
•	123320	AA496792	Hs.139572	EST	0.098
	129169	N31641	Hs.109058	ribosomal protein S6 kinase; 90kD; polypeptide 5	0.098
65	135093	U51333	Hs.159237	hexokinase 3 (white cell)	0.098
	113269	T65159	Hs.85044	ESTs	0.098
	124283	H86783	Hs.194136	ESTs; Moderately similar to zinc finger protein RIN ZF [R.norvegicus]	0.098
	114376	GMCSF	,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	Accession not listed in Genbank	0.099
	100881	HG4458-HT4727		Immunoglobulin Heavy Chain, Vdjc Regions (Gb:L23563)	0.099
	100001	1191100 1117/6/			5.556

	116572	D45654	Hs.65582	DKFZP586C1324 protein	0.099
	123956	AA621747	Hs.112847	EST	0.099
	100818	HG4018-HT4288		Opioid-Binding Cell Adhesion Molecule	0.099
	132754	W47419	Hs.56007	Human DNA from chromosome 19-specific cosmid F25965; genomic sequence	0.099
5	112741	R93080	Hs.35035	ESTs	0.099
•	112748	R93299	Hs.166492	ESTs	0.099
•	130858	S57235	Hs.246381	CD68 antigen	0.099
	124870	R69233	Hs.101504	ESTs	0.099
	125304	Z39833	Hs.124940	GTP-binding protein	0.099
10	121297	AA401995	Hs.97860	ESTs	0.099
	128602	AA046103	Hs.102367	ESTs	0.099
	124062	H00440	Hs.144524	ESTs; Weakly similar to signal transducer and activator of	
				transcription 2 [M.musculus]	0.099
	100547	HG2149-HT2219		Mucin (Gb:M57417)	0.099
15	105652	AA282505	Hs.19015	ESTs	0.099
	133390	AA459945	Hs.72660	KIAA0585 protein	0.099
	133503	M33195	Hs.743	Fc fragment of IgE; high affinity I; receptor for, gamma polypeptide	0.099
	109461	AA232667	Hs.58210	ESTs	0.099
	102068	U09117	Hs.80776	phospholipase C; delta 1	0.099
20	113464	T86931	Hs.16295	ESTs	0.099
	104240	AB002368	Hs.70500	KIAA0370 protein	0.099
	121113	AA399109	Hs.161813	ESTs	0.1
	122896	AA469952	Hs.97899	ESTs; Weakly similar to dal2; len:343; CAI: 0.17; ALC_YEAST P25335	
				ALLANTOICASE [S.cerevisiae]	0.1
25	102405	U43148	Hs.159526	patched (Drosophila) homolog	0.1
	103599	Z33905	Hs.81218	receptor-associated protein of the synapse; 43kD	0.1
	121079	AA398719	Hs.14169	ESTs; Weakly similar to CREB-binding protein [H.sapiens]	0.1
	115820	AA427487	Hs.39619	ESTs; Weakly similar to RETICULOCALBIN 1 PRECURSOR [H.sapiens]	0.781
	125106	T95766	Hs.189760	ESTs	0.1
30 .	131373	N68116	Hs.26146	Down syndrome critical region gene 3	0.1
	120224	Z41239	Hs.106960	ESTs	0.1
	133090	AA448228	Hs.6468	ESTs	0.1
	132300	AA133244	Hs.44234	ESTs	0.1
0.5	113129	T49384	Hs.8988	EST	0.1
35	110638	H73197	Hs.17241	ESTs	0.1
	131364	R53255	Hs.26010	ESTs	. 0.1
	105370	AA236476	Hs.22791	ESTs; Weakly similar to transmembrane protein with EGF-like and two	0.000
				follistatin-like domains 1 [H.saplens]	0.238

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TABLE 11A shows the accession numbers for those primekeys lacking unigeneID's for Table 11. For each probeset we have listed the gene cluster number from which the oligonucleotides were designed. Gene clusters were compiled using sequences derived from Genbank ESTs and mRNAs. These sequences were clustered based on sequence similarity using Clustering and Alignment Tools (DoubleTwist, Oakland California). The Genbank accession numbers for sequences comprising each cluster are listed in the "Accession" column.

Pkøy: CAT number: Accession:	Unique Eos probeset identifier number Gene cluster number Genbank accession numbers					
Pkey CAT number	Accession					
100610 19864_1	AW161357 Al879062 Al928938 AW161097 AW161167 BE314465 AA351715 F0709£ AA179034 F08510 F00653 Al936671 AA476718 AW772454 Al807703 R44253 AA976667 Al985186 Al650254 H38942 R84829 AA018724 AA001000 H85934					
	AA019126 H8509 AA017000 AA339355 AW950556 D51397 AA213981 BE548002 AI056359 AA001560 AW952113 AA317769 AI857477 AI857745 AW249771 AW162661 H38943 AA018628 R85885 AI984613 AI934765 AI796172 AW157488 AI929191 R85523 D51221 D53851 H85510 AI749674 F21582 AA323145 AA019127 AA687444 T06745 AI699293 H29532 AA214029 AA223656 NM_016834 X14474 R19697 H09695 R17455 R13812 R19056 AI681231 AI590200 R37671 AA861828 AI990023 AI935669 AW005821 AA324581 H17335 R37659 R42802 R46242 R60936 R59731 H28993 AA479907 R44570					
	AI890696 AA308884 AA507078 R41274 AI365507 T16348 AI560453 F03259 F04722 T16312 AA016081 AW073061					
100674 21517_2	BE314824 W28930 R44098 R51045 AW403342 AW248986 BE561709 AA357312 BE311834 BE389496 BE294887 AW732696 BE047868 AI702383 BE019155 AI702367 BE408966 BE280458 BE313759 BE513492 BE535404 BE280258 AC005263 NM_007165 L21990 AW732711 AI564920 AW249094 BE265365 AW607186 AW607346 BE005217 H27211 U46230 BE260066 BE207043 BE546782					
•	AW248659					
108559 41469_9	AA085228 AA085161					
100721 19818_1	L40904 NM_005037 X90563 AB005526 H21596 AA088517					
100748 41861_1	X06096 X05826					
100750 15759_1	BE157260 BE157265 R48118 H43827 Z17877 AW379070 AW291778 M20605 J03253 M14206 V00568 Al860465 AW296022 M13930 AL047400 J00120 BE018476 AW675223 T26980 F06694 R22709 R24720 H22753 Al903100 Al903094 AW937823 X00364 D10493 K01904 K01906 K00535 L00058 AA410662 AW384760 AA304930 Al680985 X00198 H58025 AW998901 AV653447 N31654 AW610357 AW610369 AW862480 BE223010 AW3844172 AW384219 AW3844171 AW384218 AA298522					
	BE140421 AW945162 AW751711 AA514409 AW747912 Al214214 W87741 AA972406 AA554513 BE302087 Al249030 AA477850 AV653129 Al281360 Al274110 W87861 AA641366 X66258 Al051600 AA877139 AA527483 AA857219 Al250782 AA625531 AA807892 Al278811 Al224033 H24033 AA593396 AW129709 R45453 N22772 AA235530 T29737 Al016409 Al688907 AA568370 AA722760 Al539329 AA550843 AW674698 Al538452 Al538453 Al337957 AA477744 AA464600 Al140319 AW949294 Al339781 Al828736 AA923634 AA344094 Al278350 AA975567 AA908416 AA857170 AW023520					
100751 24700_1	R43413 R48004 F02958 Al989439 R11207 AA737307 D10493 AW950652 Al093842 Al474024 AA703369 R11264 M13930 M13937 Al18363 M13933 M1393 M1393 M1393 M13933 M1393 M13933 M1393 M1393 M1393 M1393 M1393 M1393 M1393 M1393 M1393 M1					
	Al190013 Al206355 Al188721 Al189429 Al189364 Al186330 Al431595 Al189595 Al188781 Al148647 Al200022 Al221552 Al220923 Al188728 AA233034 Al189807 Al189641 Al219044 Al148774 Al200658 W71989 Al207360 Al188824 Al200559 Al200270 AA644163 Al199943 Al151301 Al189555 Al262724 Al148590 Al148695 Al126906 Al149163 K03183 K03189 Al189842 Al221014 N30608 Al186465 Al220865 Al188498 Al138226 Al189968 Al221019 Al138197 Al149426 Al148904 Al186218 Al188348 Al160579 Al198460 Al149039 Al160936 Al219055 Al184784 Al221580 Al161082 Al160814 Al123896					
	Al417614 Al126101 Al188872 Al149571 Al168533 Al149072 Al149467 Al131286 N30684 Al160705 Al160692 Al149559 Al273580 Al189442 Al138448 Al149591 N27302 AA400910 Al138431 Al138435 Al128407 N30216 Al128296 Al219589 Al188492 Al149447 Al168482 H95374 Al219009 N31616 Al276216 N32233 Al291937 N30741 Al188689 N27111 R23214 Al221605 Al184348 Al200375 H94451 N26397 Al871881 AA232905 N30833 Al220780 H94446 N30822 H87464 R68815 N30290 Al128424 H12587 T47334 H87631 H87156 Al219133 Al868741 AA330859 H86993 AA330413 H93656 N30817					
,	T90191 H93668 Al200054 H95207 T47316 H95381 T49170 R00880 T49171 N27381 H94107 R63352 T85053 AW451899 H95142 N30313 H94015 H86987 T28278 N29701 C18834 AA331267 AA330939 Al654493 N27073 N29831 R68113 N30758 R26086 N32108 H95135 AA330414 AA330978 Al219422 Al189453 Al199951 X00264 NM_000894 AA371909 AA063496 T29543 AA371971 AA372026 AA371978 AA371346 Al051683 Al186418 Al220659 Al189068 Al219266 Al186552 Al188715					
100760 1334_7 100775 18179_3	Al149156 AW794626 M27126 M27014 J05581 M61170 T27692 M34088 M34089 AW860335 AW579047 AW610437 AW610386 AW610422 AW610473 AW579078 AW604897 AW860163 AW579067 AW862410 Al816584 AW177757 AW602769 Al909790 AW860331 Al909787 Al909811					

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## ASS2469 A 1917/26 A 191	5		AW001137 AI567905 T84561 AI631295 AA151351 H02932 AI884519 AA367457 AW369421 AI676846 AW391803 AI610869 AW192838 AI922289 AI952140 AI910233 AI479474 AW001395 AA488073 AI985760 AW130017 AI858369 AA627845 AW081805 AA158865 AI624443 AA344985 AA569793 R72486 AI589329 AI903204 AI269893 AA641284 AI279932 AA149270 AI697120 AA729146 AI589353 AA480067 AI923310 AA530908 AI275395 AA425062 AA580280 AA889527 AA158866 AW131341 AA573028 AA877326 T29335 AW951288 H04235 AA099243 AA994659 AI659618 AA887919 AI299297 AW001116 AW263844 AI270578 AA970828 AW572126 AA775299 AW369449 AW369398 AW369452 AI933677 AI870710
### ASSSSSS FIGURES AMM-TISSED HIZESS HIZESS TISSED ASSISTS TAM/SISSES AMMSSSSD HIGESS FIGURE DASSES FIGURE VALVES AND STATES AMMSSSSD AMMSSSSD FIGURE AMSSSSD AMSSSSSD FIGURE AMSSSSD AMSSSSD FIGURE AMSSSSD AMSSSSD AMSSSSD FIGURE AMSSSSD AMSSSSD AMSSSSD FIGURE AMSSSSD AMSSSSD AMSSSSD AMSSSSD FIGURE AMSSSSD AMS	10	100800 24735_1	AA932409 A1187328 A1672970 A1886098 AW440471 AW138860 A1866858 A1802528 A1926172 AW243914 A1933690 AA996114 AA536189 AW009937 A1918060 A1270379 A1973169 AW175638 AW369413 NM_006227 L26232 R50649 AU077024 AL008726 AA411079 R35151 BE278153 BE278139 A1459777 R88036 Z43210 F07326 AF052157 R17844 BE615476 T82160 R71985 H21963 AA299158 AW368246 R48123 R50628 R70441 H27245 H72015 R72345 R39392 A1909738 BE612778 BE613234 D52116 D52136 D52132 D52067 D51922 D51995 D51905 N34249
Tubbel 1 Tyrop 149940 Aloga2477 AM/37045 Ne2294 Al758689 AA295526 AA284586 ALS9777 AA460277 AB325559	15		AA358363 R50262 AW473860 H52335 H43953 H21964 T39505 Al887517 AW156925 AW839850 H02628 AW007705 Al561008 F22392 R71279 AA995433 R50725 W24462 R71931 AA464437 AW591731 R25667 R52695 R50810 Al560805 Al089266 H68386 H41353 H28590 AW001860 Al141623 AA250773 Al284778 AW511412 AW083975 AA130377 AW026047 R50551 R81494 Al357668 Al078272 F32666 F36981 AW304865 H43906 AA931068 R48010 Al540217 Al017339 Al291812
100816 18004_3 1700816 18004_3 1700816 18004_3 1700816 18004_3 1700816 18004_3 1700816 18004_3 1700816 18004_3 1700816 18004_3 1700816 18004_3 1700816 18004_3 1700816 18004_3 1700816 18004_3 1700816 18004_3 18008	20		T40641 T47608 T48940 Al082477 AW470145 N92284 Al758958 AA298512 AA284586 Al597777 AA480277 Al932559 Al869081 AA476615 AA503651 Al656024 AW168522 Al682051 Al689106 Al274592 Al520917 BE258916 BE615861 BE280282 R53386 BE278255 BE278398 T47607 AA477662 H68385 100817 19648_1 L34355 L46810 NM_000023 U08895 AA424260 Al097272 AA424162 N79764 F19290 F25278 Al479385
BE269598 BE559595 BE359881 BE56031 BE514199 BE56037 BE560454	25	100818 19604_3	AA453282 U79251 AA843851 R38201 R66461 R44908 AA683289 H17477 R37364 R52832 AW298336 AA351391 NM_002545 L34774
## 100898 8542 1		100881 458_127	
AW338647 AW374322 AA34040 BE293488 AL 138361 BE560260 AT745075 AA317380 AW948382 AB33411 AB63582 AB31042 AB361878 AA618606 AA729052 AI424959 AA199716 AW769374 AB28472 AW043307 AI862816 AI203583 AW493912 AA292744 AW471439 AI473989 AA593336 AA464070 AI678937 AW059451 AA970763 AA610480 AA593328 AA464090 AA768985 AI209820 AA346900 AA464718 AA99361 D61482 D55935 AI3069931 AA70695 AI600830 AA690827 AB66259 AI61928 AB51868 AI208320 AA346900 AA768985 AW158609 AW518663 BE045525 Z41251 AI868091 AA970810 AI60887 AB66259 AI619282 AB215437 AI565416 AA463934 AW168609 AW518663 BE045525 Z41251 AI868091 AA908160 AI026897 AB66259 AI619282 AB215437 AI565416 AB463934 AW168609 AW518663 BE045525 Z41251 AI868091 AA908160 AI026897 AB66259 AI619282 AB215437 AI56541 BE541097 BE255652 BE265878 BE394102 W27502 AB66259 AI619282 AB215437 AI65651 AB63370 AB66259 AI619282 AB215437 AB2647 BE3950 BE39540 AI036606 AIL036420 U35630 AW298574 AB66259 AI619282 AB3659 AB3758 AB3155 F18090 AB5755 BE3956 BE3957 BE3957 BE3958	30		X07881 NM_006249 X07637 AA376715 AA376677 X07715 X07704 S80916 BE387614 R51501 AA199714 AW674779 F08178 BE269071 AA376313 H08264 AA380420 H18785 AL042151 BE277758 BE267438 NM_005850 L35013 BE540833 BE390902 BE391494 BE277459 BE385592 BE390612 BE384263 BE387779
40 102459 3556_1 102620 16821_37 A976427 U66052 45 102673 24986_6 10275 5145_4 102753 2226_1 102793 3462_4 102896 108898 NA93668 108266 575578 AA425061 AA429317 AI816143 AA910669 AI286022 AI286019 103522 21640_1 103522 21640_1 103522 21640_1 103522 21640_1 103523 21640_1 103	35		AW938647 AW374332 AA434040 BE293488 AL138361 BE560260 AI745075 AA317980 AW949382 AI834311 AI653582 AI831042 AI361878 AA618606 AA729052 AI424969 AA199715 AW769374 AI828422 AW044307 AI862816 AI203583 AW084461 AW514655 AA831883 AA290672 AA831286 AA578510 AW089965 AW150746 AA292743 H22232 AI469275 AW439312 AA292744 AW471443 AI473989 AA593336 AA464070 AI678937 AW069451 AA970763 AA610480 AA593328
45 102673 2498-6 102675 5145_4 172509 172512 798357 R31335 F18090 172512 798357 R31335 F18090 172512 798357 R31335 F18090 18262 R35578 AA425061 AA429317 AI815143 AA910669 AI286022 AI286019 102799 34624_4 127034 51148_2 182961 NN_00663 U80226 S75578 AA425061 AA429317 AI815143 AA910669 AI286022 AI286019 1829034 51148_2 182961 NN_00663 U80226 S75578 AA425061 AA429317 AI815143 AA910669 AI286022 AI286019 1829034 51148_2 182961 NN_00663 U80226 S75578 AA425061 AA429317 AI815143 AA910669 AI286022 AI286019 1829034 AI815175 182903 AI815175 BE322229 AA296468 AW959763 BE276164 BE264109 1829229 AI815175 BE326229 AI829276 AA427485 AA496400 AA352389 1829664 N63044 N51226 AI5151248 AI521996 AI924777 AW375954 AI860275 W00549 AI742673 AW612288 AI763062 AA632510 AI087347 AI088070 AI214349 AA890297 AI494156 AI698598 AA631658 AA504593 AA860733 AI266761 AW663214 AW771231 AA639610 AI769806 AI769746 AW014326 AI288611 AA429212 W00881 119388 1762256_1 AA429212 W00881 126856 291965_1 AA629212 W00881 AA696567 AA488977 AA908902 AI637637 AA927056 AI870139 AW340492 AA488755 AA129794 AA306523 AA554253 BE256277 AC053467 AW962084 A366523 AA54253 BE256277 AC053467 AW962084 AA9066248 R34447 AA847136 AI684489 AI523112 AW044269 AI379138 N29386 AA761543 N79248 AA960845 AA768316 AI147926 AI718599 AI880620 R67457 AI216016 AI738663 H04648 NW024868 AI133339 AI859206 R51175 U87167 BE379324 BE392008 AA340819 AA343110 T57275 D59164 AW2299312 AI43442 AI83339 AW024975 R40262 AW269126 R09430 T566500 AI367247 AI255132 BE464248 T58658 AW207785 T58607 R51194 AI732276 R53587 AI820697 AW26498 AW269126 R09430 T566500 AI367247 AI255132 BE464248 T58658 AW207785 T58607 R51194 AI732276 R53587 AI820697 AW060526 BE550034 W30689 AW271859 AA411456 AI341551 AA242990 AA243027 H87046 D20360 AI184053 AA146956 AA600526 BE550034 W30689 AW271859 AA411456 AI341551 AA242990 AA243027 H87046 D20360 AI184053 AA146956	40	126126 1630017_1	Al886259 Al612932 AA215437 Al956014 BE541087 BE255652 BE265878 BE394102 W27502 U48936 L36592 X87160 NM_001039 AL036606 AL036420 U35630 AW298574 W80551 M85370
BE392626 BE256735 AA301453 N55872 H01676 AA292746 AA427485 AA496400 AA352389	45	102673 24986_6 102675 5145_4 102753 2226_1 102799 34624_4	Al457548 U72509 U72512 T98357 R31335 F18090 L32961 NM_000663 U80226 S75578 AA425061 AA429317 Al815143 AA910669 Al286022 Al286019 U88896 U88898 AA916056 T03285 Al341594 Al359534 Al634031 U88897
AA639610 Al769806 Al769746 AW014326 Al288611 127071 188097_1	50	_	BE392626 BE256735 AA301453 N55872 H01676 AA292746 AA427485 AA496400 AA352389 Y10518 Y10514 Z83935 Y10508 AK000055 Y10519 AI142012 AI681175 BE222219 AA890586 BE504347 BE328064 N63044 N51226 AI151248 AI521996 AI924777 AW375954 AI860275 W00549 AI742673 AW612288 AI763062 AA632510 AI087347
126856 20669_1 Al084125 Al083773 Al479687 Al939609 Al968662 AF129507 NM_013282 AW971840 AW298508 AA744240 AA811217	55	126456 291965_1	AA639610 AI769806 AI769746 AW014326 AI288611 AA250806 AA459220 AA429212 W00881
60 103996 224545_1			AI084125 AI083773 AI479687 AI939609 AI968662 AF129507 NM_013282 AW971840 AW298508 AA744240 AA811217 AA827671 AA811055 AA806567 AA488977 AA908902 AI637637 AA927056 AI870139 AW340492 AA488755 AA129794
AW028488 Al133339 Al859205 R51175 U87167 BE379324 BE392008 AA340819 AA343110 T57275 D59164 AW299312 Al434422 Al936390 AW024975 R40262 AW269126 R09430 T56590 Al367247 Al253132 BE464248 T58658 AW207785 T58607 134947 844579_1 129311 16078_1 AK000526 BE550084 W30689 AW271859 AA411456 Al341551 AA242990 AA243027 H87046 D20360 Al184053 AA146956	60	103996 224545_1	AA321355 AW964592 R23284 H73883 R23382 N47914 C01377 H04668 AW606248 R34447 AA847136 Al684489 Al523112 AW044269 Al379138 N29366 AA761543 N79248 AA960845 AA768316 Al147926 Al718599 Al880620 R67467 Al216016 Al738663 H04648
134947 844579_1 R51194 Al732276 R53587 Al820697 129311 16078_1 AK000526 BE550084 W30689 AW271859 AA411456 Al341551 AA242990 AA243027 H87046 D20360 Al184053 AA146956	65	113213 23798_1	AW028488 A1133339 Al859205 R51175 U87167 BE378324 BE392008 AA340819 AA343110 T57275 D59164 AW299312 Al434422 Al936390 AW024975 R40262
		_	R51194 AI732276 R53587 AI820697 AK000526 BE550084 W30689 AW271859 AA411456 AI341551 AA242990 AA243027 H87046 D20360 AI184053 AA146956

			AAA31522 AA301989 AI909058 D12149 N41960 BE222214 AA609922 AA828176 AA393359 AA398693 AW024956
•			BE467805 AW298623 AW264085 Al024454 Al024719 Al431927 T55087 Al611014 T54920 AA131253 Al436344
		9724_2	AA017176 Al359979 AA047836 AA017063 AA016303 AA001545
5		110077_1 156215	AA063315 AA063316 AF015910
5	100515		AA305746 D90187 T63943 AW951154 T29182 AI734941 D13264 AI299239 Z18812 AW299859 W24476 AA933064
	100010	012_1	AA489759
	100531	46038_1	AW888554 AW607282 AA319986 M28590
4.0		22955_11	M55405 AW752552
10	100574	17320_2	AA326895 M10036 NM_000365 N84665 H69414 N84657 AA380453 AA329743 AA357367 AA188770 AA376532 AA353653
			AA158953 AA083176 BE537313 AA181433 D53373 R57376 AA206698 R14807 H18899 H11191 H93892 R25593 T61134
			N93285 AA083081 AA831789 H13137 AA497014 AA079330 AA182861 H13138 W47161 R62913 AA687089 AA211112 AA429237 AL035923 AA100070 AW392898 AI566433 AA866006 AA214002 AW392865 N79454 AA197181 AI680371
			AA176501 AA737967 AI089225 F34874 AW571437 AI620620 AA573489 AA423816 AA164917 AA458455 T47072 AI569087
15			Al261656 AA730919 Al633441 AW195182 Al351622 AW243465 Al872649 Al359227 AA987941 Al693770 T47073 AW779948
			AW510580 Al635626 AW627601 AA864326 AA953578 Al341418 BE222853 Al241963 Al094663 AA928380 AA493373
			AW043762 Al377783 AW958987 BE619760 AA385240 BE277975 BE280095 AW631443 AA581048 BE618715 BE299610
			C14874 BE559858 BE378455 BE618290 BE544585 Al525575 BE548897 BE267110 AA804738 BE269821 AA918133
20			BE277647 AA599947 BE280735 BE390239 N74150 T12504 AI208197 AW955527 AA113897 N40081 H73835 H70393 AI434041 W22950 AI192661 BE264461 W26486 AA626424 AA196694 T69209 AA857976 AI540287 AA410599 AA864287
20			AW950564 AA013320 T49283 AI541438 AW804703 AA335534 AA335659 BE562269 BE618802 BE277850 BE546413
			BE280994 AA204813 BE561694 BE543524 BE253647 AW001452 W19116 BE542508 AA205894 BE254675 BE270033
			Al525906 BE251792 AA975700 BE272138 AW607671 N87686 M10036 BE515060 BE298607 Al745178 U47924 H03193
05		tigr_HT2798	725424
25		tigr_HT3768	M88357
		tigr_HT3846 tigr_HT4265	L29141 M69180 M81105 L33999
		tigr_HT4383	U04688
		tigr_HT4504	U09806
30			U12139 .
		genbank_T91518	
		tigr_HT688 _entrez_W38150	X65561
		entrez_U38268	
35		entrez U51010	
		entrez_U67092	
		genbank_N74496	
			K01160
40		entrez_K03430 genbank_AA0704	
		483241_1	AA070853 AA075749 AA075716
		genbank_AA0790	
		genbank_AA1289	
45		entrez_M60299	
43	117437	entrez_M64358 genbank_N27645	M04530 N27645
	101798	entrez_M85220	M85220 ·
			S69265
50			Y10141
50			Z26256
		genbank_T54095	
		genbank_R48295 NOT_FOUND_en	
		NOT_FOUND_en	
55		tigr_HT1067	M22406
		tigr_HT2219	M57417
	100564	tigr_HT2324	Z11585

TABLE 12: shows genes, including expression sequence tags, that are down-regulated in prostate tumor tissue compared to normal prostate tissue as analyzed using Affymetrix/Eos Hu01 GeneChip array. Shown are the ratios of "average" normal prostate to "average" prostate cancer tissues.

10	Pkey: ExAccn: UnigeneID: Unigene Title:	Unique Eos probeset identifier number Exemplar Accession number, Genbank accession number Unigene number Unigene gene title
	R1:	Background subtracted normal prostate : prostate tumor tissu

1	5	Pkey	ExAcon	UnigenelD	Unigene Title	R1
		100522	HG1763-HT17	80	Prolactin-Induced Protein	17.4
		130803	M81650	Hs.1968	semenogelin I	16.785
		118068	N53943	Hs.13743	ESTs	13.225
2	0	114251	Z39898	Hs.21948	ESTs	12.7
		112134	R46025	Hs.7413	ESTs	8.735
		101436	M20642	Hs.158295	Human alkali myosin light chain 3 mRNA; complete cds	8.175
		104028	AA361094	Hs.221128	ESTs	8.15
_	_		AA149204		ESTs; Highly similar to growth arrest inducible gene product [H.sapiens]	7.535
2	5		AA174173	Hs.12622	ESTs	7.212
			AA251741	Hs.25882	DKFZP586M1824 protein	7.175
			H29231	Hs.27384	ESTs	6.701
			AA761378	Hs.192013		6.642
2	^		N35229	Hs.7037	pallid (mouse) homolog; pallidin	6.411
3	U		U48807	Hs.2359	dual specificity phosphatase 4	6.395
			W90385	Hs.15082	ESTs	6.15 6
			L33881	Hs.1904	protein kinase C; iota	6
			AA491209 T49655		ESTs; Weakly similar to reverse transcriptase [M.musculus] ESTs; Modly smir to !! ALU SUBFAMILY SQ WARNING ENTRY !! [H.sapiens]	5.95
3:	5		M61906	Hs.6241	phosphoinositide-3-kinase; regulatory subunit; polypeptide 1 (p85 alpha)	5.925
٦.	,		· AA399218	Hs.92423	ESTs	5.7
			AA281780		ESTs; Weakly similar to similar to Kruppel-like zinc finger protein [C.elegans]	5.7
			F10192		Tubulin; alpha; brain-specific	5.625
			X80507	Hs.8939	yes-associated protein 65 kDa	5.5
4	0		AA234048	Hs.7753	calumenin	5.486
•	•		AA233122		ESTs; Highly similar to multifunctional calcium/calmodulin-dependent protein	
					kinase II delta2 isoform [H.sapiens]	5.4
		131699	R68657	Hs.90421	ESTs; Modly smlr to !! ALU SUBFAMILY SX WARNING ENTRY !! [H.sapiens]	5.279
	_	104490	N71503	Hs.43087	ESTs; Weakly similar to dysferlin [H.sapiens]	5.266 .
4:	5	102124	U14528	Hs.29981	solute carrier family 26 (sulfate transporter); member 2	5.151
		109280	AA196635	Hs.86081	ESTs	5.134
			F09739		Homo sapiens mRNA full length insert cDNA clone EUROIMAGE 21920	5.075
			AA045709	Hs.40545	ESTs	5.075
_	^		M21665	Hs.929	myosin; heavy polypeptide 7; cardiac muscle; beta	5.055
5	U		R80664	Hs.77067	ESTs	5.033
			R62444		KIAA0931 protein	4.675
			M20543	Hs.1288	actin; alpha 1; skeletal muscle	4.626
			R93943	11-004404	yt72c12.r1 Soares retina N2b4HR Homo sapiens cDNA clone IMAGE:275735 5',	4.6 4.559
5:	<		W87544	Hs.221184		4.451
J.	J		J03460 R56068	Hs.99949 Hs.4268	prolactin-induced protein ESTs	4.45
				Hs.53106	TILL	4.45
			R16833 R40873		KIAA0432 gene product	4.301
			AA448286		ESTs; Highly similar to atrophin-1 interacting protein 4 [H.sapiens]	4.2
6	n		AA447006	115.30074	ESTs; Moderately similar to II ALU SUBFAMILY SQ WARNING	4.175
0.	•		N32787	Hs 11123	ESTs; Moderately similar to hypothetical protein 2 [H.sapiens]	4.1
			Z84483		Human DNA sequence from PAC 46H23, BRCA2 gene region chromosome 13q1:	
			Z40186	Hs.26409		4.05
			AA262491	Hs.186572		4.048
6	5		AA609749		ESTs; Highly similar to unknown protein [R.norvegicus]	4.041
			H97993		ESTs; Weakly similar to KIAA0512 protein [H.sapiens]	4.028

	120217	Z41078	Hs.66035	ESTs	4.028
		AA084524		zn19d8.s1 Stratagene neuroepithelium NT2RAMI 937234 Homo sapiens cDNA	4.023
		AA400030	Hs.8360	ESTs; Weakly similar to II ALU CLASS B WARNING ENTRY II [H.sapiens]	3.925
	120418	AA236010	Hs.26613	Homo sapiens mRNA; cDNA DKFZp586F1323 (from clone DKFZp586F1323)	3.91
5		N74897		DEAD/H (Asp-Glu-Ala-Asp/His) box polypeptide 15	3.889
		T80174		ESTs; Moderately similar to similar to NEDD-4 [H.sapiens]	3.85
		AA099585		ESTS	3.833
		X61100	Hs.8248	NADH dehydrogenase (ubiquinone) Fe-S protein 1 (75kD) (NADH-coenzyme	3.818 3.792
10		H06773 AA258144	Hs.93850 Hs.221576	protein kinase; AMP-activated; gamma 2 non-catalytic subunit	3.779
10		R37460	Hs.25231		3.768
		W86600	Hs.9842	ESTs	3.75
		AA481254	Hs.30120		3.708
	129775	R94659	Hs.12420		3.707
15		H20568		phospholipase A2-activating protein	3.7
		R87160	Hs.33665		3.7
		AA375791	Hs.131894		3.674 3.653
		W92797		DKFZP434G162 protein dachshund (Drosophila) homolog	3.625
20		AA252079 AA242751		KIAA0903 protein	3.62
20.		AA487228		ESTs	3.614
		AA024664		NADH dehydrogenase (ubiquinone) 1 alpha subcomplex; 5 (13kD; B13)	3.613
	124780	R42493	Hs.220839		3.6
	130631	AA025399	Hs.169737		3.592
25		AA211320	Hs.79404	neuron-specific protein	3.568
	104160	AA455706		ESTs; Weakly similar to 78 KD GLUCOSE REGULATED PROTEIN	3 550
	400004	4.4050450		PRECURSOR	3.559 3.542
		AA258158	Hs.22153 Hs.176586	ESTs; Weakly similar to KIAA0352 [H.sapiens]	3.525
30		H19673 AA233299	Hs.72158		3.522
50		F02367	Hs.27252		3.5
		AA257107	Hs.194331	ESTs	3.5
		AA455653	Hs.44581	ESTs; Weakly similar to HEAT SHOCK 70 KD PROTEIN 6 [H.sapiens]	3.459
	120524	AA261852	Hs.192905	ESTs	3.45
35		H74330	Hs.150000		3.425
		AA256976	Hs.18800	EST's; Weakly similar to KIAA0579 protein [H.sapiens]	3.42
		X05451	Hs.158295 Hs.49829	Human alkali myosin light chain 3 mRNA; complete cds	3.417 3.407
		N70298 Al028384	Hs.127331		3.399
40		AA159953		ESTs; Weakly similar to anyisulfatase B precursor [H.sapiens]	3.325
		AA600116	Hs.112526		3.318
		N50866	Hs.47135		3.317
		AA287097		transcription factor 4	3.315
45		H85897	Hs.27755		3.309
45		AA342104	Hs.96777		3.3 3.295
		AA278824 AA946876	Hs.19218 Hs.148376		3.292
		HG4020-HT4		Transglutaminase	3.288
		D29956		ubiquitin specific protease 8	3.273
50		AA608903	Hs.106220	KIAA0336 gene product	3.269
	134719	L07515	Hs.89232	chromobox homolog 5 (Drosophila HP1 alpha)	3.266
		H29209		ESTs; Highly similar to FYVE finger-containing phosphoinositide kinase [M.muscu	lus] 3.26
		AA418880	Hs.185797	ESTS	3.212 3.197
55		R60523	Hs.109087		3.179
23		AA970504 R94500	Hs.146103 Hs.108046		3.175
		AA448164	Hs 99153	ESTs; Highly similar to CGI-73 protein [H.sapiens]	3.151
		AA431302	Hs.98721	EST; Weakly similar to N-copine [H.sapiens]	3.151
		X85134		retinoblastoma-binding protein 5	3.15
60	129937	M95767	Hs.135578	chitobiase; di-N-acetyl-	3.15
		AA057341		helicase-moi	3.15
		AA018219	Hs.226923		3.125
		AA421773	Hs.161008		3.125 3.12
65		AA149007		Ets homologous factor	3.11
05		N48818 AA485973	Hs.46884 Hs.143947		3.104
		AA400080	Hs.97774	ESTs	3.1
		T80620	Hs.186473		3.075
		AA401739	Hs.5111	ESTs	3.066

	119767	W72562	Hs.58119	ESTs	3.057
	115776	AA424038	Hs.58197	ESTs	3.056
		R22988	Hs.220950	ESTs	3.05
	115301	AA280047	Hs.43948	ESTs	3.05
5	118448	N66412	Hs.49189	ESTs	3
	106586	AA456598	Hs.256269	ESTs	2.995
	110415	H48239	Hs.29739	ESTs; Weakly similar to RAS-RELATED PROTEIN RAB-3A [H.sapiens]	2.979
	105173	AA182030	Hs.8364	ESTs	2.978
		L07594	Hs.79059	transforming growth factor; beta receptor III (betaglycan; 300kD)	2.976
10		H58383	Hs.258544		2.976
		R24464		KIAA1102 protein	2.964
		HG4058-HT43		Oncogene Aml1-Evi-1, Fusion Activated	2.957
		AA481068	Hs.31835	ESTs	2.95
		D11930	Hs.3592	ESTs	2.95
15		N68869	Hs.15119		2.936
13		T93795	Hs.17252		2.917
		AA210695	Hs.17212		2.917
		AA490227	Hs.105252		2.904
		W87465		ESTs; Weakly similar to D2092.2 [C.elegans]	2.9
20		AA465160	Hs.63386		2.9
20					2.896
		AA432243	Hs.41086 Hs.113716		2.895
		R22905			2.895
		W78107		ESTs; Weakly similar to Yer140wp [S.cerevisiae]	2.894
25		N64378		ESTs; Highly similar to unknown function [H.sapiens]	
25		AA280300	Hs.191346		2.886
		AA152263		KIAA0849 protein	2.883
		H03686		Ras-GTPase-activating protein SH3-domain-binding protein	2.879
		W73788	Hs.43213		2.875
20		R59881	Hs.7503	ESTs	2.871
30		AA028171	Hs.153688		2.868
		Al088155		ESTs; Weakly similar to unknown [H.sapiens]	2.866
		AA280738	Hs.128679		2.863
		AA482019	Hs.238202		2.86
25		D84239		IgG Fc binding protein	2.856
35		W45491	Hs.172609	nucleobindin 1	2.854
		W01626		za36e07.r1 Soares fetal liver spleen 1NFLS Homo sapiens cDNA clone	2.852
		AA227972	Hs.43635	ESTs	2.85
		AA142919	Hs.5558	ESTs	2.847
40		R34531		KIAA0480 gene product	2.846
40		AA398720	Hs.177953		2.838
		R73816	Hs.17385		2.836
		R26206		KIAA0701 protein	2.825
		AA427783	Hs.77910	3-hydroxy-3-methylglutaryl-Coenzyme A synthase 1 (soluble)	2.816
	116355	AA504356	Hs.88650	ESTs	2.813
45	115316	AA280627	Hs.57846	ESTs	2.806
	129677	U48736	Hs.198891	serine/threonine-protein kinase PRP4 homolog	2.8
	130971	H20332	Hs.28707	signal sequence receptor; gamma (translocon-associated protein gamma)	2.799
	115054	AA252863	Hs.87729	ESTs	2.795
	130285	AA063546	Hs.202968		2.792
50		H93575		Homo sapiens mRNA; cDNA DKFZp564J142 (from clone DKFZp564J142)	2.783
	125502	AA732329	Hs.191959	ESTs	2.778
	114800	AA159825	Hs.131887	ESTs; Weakly similar to ORF YNL227c [S.cerevisiae]	2.768
	128625	AA242816	Hs.102652	ESTs; Weakly similar to KIAA0437 [H.sapiens]	2.766
	130159	H51098	Hs.151310	PDZ domain protein (Drosophila inaD-like)	2.75
55	107127	AA620504	Hs.22119		2.742
	113547	T90746	Hs.15233	ESTs	2.734
		AA004622	Hs.18214	ESTs	2.727
	127609	AA622559	Hs.150318	ESTs	2.726
	106922	AA490964	Hs.10056	ESTs	2.725
60		R52088		yg85c3.s1 Soares infant brain 1NiB Homo sapiens cDNA clone	2.725
	124333	H98683	Hs.154054		2.708
		N36421		ESTs; Weakly similar to SODIUM- AND CHLORIDE-DEPENDENTGLYCINE	
			TRANSP		2.706
	101609	M54927	Hs.1787	proteolipid protein 1 (Pelizaeus-Merzbacher disease; spastic paraplegia 2;	
65			uncomplica		. 2.704
	117142	H96908	Hs.42251	_ ·	2.7
		R79147	Hs.203365		2.695
		AA481505	Hs.13797		2.68
		N25996	Hs.179833		2.675
	12-7011				

	101026	J04970		carboxypeptidase M	2.675
		N66393 `	Hs.102754	•• •	2.675
		H02494	Hs.101615		2.671
		R12777.		ESTs; Weakly similar to ARGINYL-TRNA SYNTHETASE [H.sapiens]	2.66
5		N49602	Hs.13308		2.65
,		N54839		ESTs; Highly similar to mediator (H.sapiens)	2.633
					2.63
		AA421714		KIAA0896 protein	
		U32519		Ras-GTPase-activating protein SH3-domain-binding protein	2.626
10		R62452	Hs.191265		2.625
10		AA451679	Hs.194410		2.625
		X90568	Hs.172004		2.624
		N63391	Hs.9238	ESTs	2.619
		AA459242		ESTs; Weakly similar to Kelch motif containing protein [H.sapiens]	2.615
	112079	R44164	Hs.23014		2.6
15	123033	AA481271	Hs.193945	ESTs	2.591
	124196	H52617	Hs.144167	ESTs	2.586
	125873	H14437		yl25a04.r1 Soares breast 3NbHBst Homo sapiens cDNA clone	2.58
	117684	N40184	Hs.45050	ESTs	2.575
	134938	D30037	Hs.168326	phosphotidylinositol transfer protein; beta	2.575
20	131822	AA215647	Hs.200332	ESTs	2.568
	135185	U71203	Hs.96038	Ric (Drosophila)-like; expressed in many tissues	2.564
		N40467	Hs.93834	FSTe	2.557
		N78582		protein kinase; AMP-activated; beta 2 non-catalytic subunit	2.552
		AA405657	Hs.128791	Human DNA sequence from clone 967N21 on chromosome 20p12.3-13. Contains	
25		AA235112		ESTs; Moderately similar to similar to murine RNA-binding protein [H.sapiens]	2.549
		AA426017		ESTs; Highly similar to DNA-REPAIR PROTEIN COMPLEMENTING	2.548
		H20560	Hs.244624		2.548
		AA045333		ESTs; Weakly similar to !! ALU SUBFAMILY SB2 WARNING ENTRY !! [H.sapiens]	
		F09729	Hs.12780		2.537
30		AA410391	Hs.94592		2,525
30		R37027	Hs.5737	KIAA0475 gene product	2.525
		R73588	Hs.101533		2.525
		AA174190	Hs.50932		2.525
		R22196	Hs.34492		2.519
35		AA234362			2.507
JJ					2.5
		T23555	Hs.103288 Hs.99601		2.499
		AA496258			2.491
		AA426270			2.491
40		AA496884	Hs.23972		2.483
40		AA018042	Hs.95078		
		U60808			2.481
		X56741	Hs.5947		2.475
		R38100	Hs.106294		2.475
45		T15665	Hs.6185		2.475
43		W58202	Hs.125731		2.475
		R62447	Hs.22123		2.471
		AA600323	Hs.112535		2.462
		U95020			2.457
50		U42390			2.455
50		T56056	Hs.9992		2.452
		L11066	Hs.3069		2.448
		AA280583	Hs.256501		2.437
	128211	Al206427	Hs.166707		2.43
		L11005	Hs.81047		2.425
55	115985	AA447709	Hs.132094		2.425
		N90041	Hs.9585		2.418
	129430	AA258842			2.418
	133863	C13990	Hs.76930	synuclein; alpha (non A4 component of amyloid precursor)	2.417
	111164	N66857			2.416
60	132143	AA257056	Hs.7972	KIAA0871 protein	2.412
	130330	M55047	Hs.154679	synaptotagmin 1	2.408
		Z39451	Hs.27389		2.406
		H94043	Hs.24341		2.403
		AA034325	Hs.54320		2.4
65		N62506	Hs.21958		2.4
J J		AA195405	Hs 110347	et : :	2.397
		N69998	Hs.21801		2.394
		AA608955	Hs.109653		2.389
		AA608893	Hs.190065		2.388
	140004	~~~~~	113.130003	24.4	

		AA262821	Hs.28578		2.385
	103143	X66141	Hs.75535		2.384
	123645	AA609310	Hs.188691	ESTs	2.383
	130123	AA001835		zinc finger protein 262	2.379
5	-	AA428368		ESTs	2.378
•		AA436666	Hs.59761	ESTS	
					2.375
		W45574	Hs.252497		2.372
	123973		Hs.182151		2.361
	135197	U76456		Homo sapiens tissue inhibitor of metalloproteinase 4 mRNA, complete cds	2.357
10	118689	N71545	Hs.184544	ESTs	2.357
	107734	AA016225	Hs.93386	ESTs	2.354
	124590		Hs.41381	ESTs; Weakly similar to ubiquitin hydrolyzing enzyme I [H.sapiens]	2.35
	111163			ESTs	
			Hs.17606	## · *	2.348
1.5	112349		Hs.22665	ESTs; Moderately similar to dJ83L6.1 [H.sapiens]	2.345
15		AA262179	Hs.169343		2.345
	134238	R81509	Hs.184571	splicing factor; arginine/serine-rich 11	2.341
	116766	H13260	Hs.95097		2.336
	106331	AA436853	Hs.34795	ESTs	2.333
		AA443752	Hs.10784		2.332
20		AA599814	Hs.46637		2.332
20	124697			ECTs: Mediu amiste II At II CHIDEANII V. I MADAINO ENTRY II II Carisma	
				ESTs; Modly smir to !! ALU SUBFAMILY J WARNING ENTRY !! [H.sapiens]	2.322
		AA176688	Hs.221139		2.313
		AA304993		ESTs; Weakly similar to p60 katanin [H.sapiens]	2.307
~ ~		AA252621	Hs.93842	ESTs	2.301
25	119819	W74371	Hs.58383	ESTs	2.297
	102302	U33052	Hs.69171	protein kinase C-like 2	2.288
	130596		Hs.16475	ESTs	2.282
	114161		Hs.22385	ESTs; Weakly similar to KIAA0970 protein [H.sapiens]	2.278
	130542		,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	Human sperm membrane protein BS-63 mRNA, complete cds	
30			He googg		2.277
20	104491		Hs.39328	ESTs	2.275
	116988			ys69e12.s1 Soares retina N2b4HR Homo sapiens cDNA clone	2.275
		AA370120	Hs.7870	ESTs; Weakly similar to YIr350wp [S.cerevisiae]	2.273
	108800	AA129731	Hs.90424	ESTs	2.273
~ =	101310	L41607	Hs.934	glucosaminyi (N-acetyl) transferase 2; I-branching enzyme	2.269
35	126842	W19498	Hs.21085	ESTs	2.255
	127251	AA936428	Hs.128638	ESTs	2.251
	124647	N91947	Hs.125033	ESTs	2.249
	127112	Al143906	Hs.125103	ESTs	2.247
	101973			UDP-N-acetyl-alpha-D-galactosamine:polypeptide	2.246
40		AA398302	Hs.127437		2.245
		AA599583		HMBA-inducible	2.243
	119980				
				heterogeneous nuclear protein similar to rat helix destabilizing protein	2.243
	124222		Hs.222844		2.24
15	129199		Hs.128629		2.236
45		AA479101		ESTs; Weakly similar to !! ALU SUBFAMILY SQ WARNING ENTRY !! [H.sapiens]	2.231
	126160	N90960	Hs.247277	ESTs; Weakly similar to transformation-related protein [H.sapiens]	2.229
	104627	AA001976		ESTs	2.228
	106474	AA450212	Hs.42484	Homo sapiens mRNA; cDNA DKFZp564C053 (from clone DKFZp564C053)	2.226
	113096	T40927		ESTs	2.225
50		AA452822	Hs.99027		2.225
-	135344			ESTs; Moderately similar to TRF1-interacting ankyrin-related	
					2.225
		AA508354		ESTs; Moderately similar to AKT3 protein kinase [H.sapiens]	2.222
		AA397841	HS.180141	cofilin 2 (musde)	2.218
ے ہے		AA026385		ESTs; Moderately similar to II ALU SUBFAMILY SB2 WARNING	2.217
55		AA033562	Hs.151572		2.212
	109292	AA199828	Hs.188662	ESTs	2.212
	104257	AF006265	Hs.9222	estrogen receptor-binding fragment-associated gene 9	2.209
	132932	T15482		ESTs	2.204
		AA262728		Homo sapiens clone 24590 mRNA sequence	2.204
60		AA004652		ESTs	2.2
		AA449828	Hs.99229		2.195
	124307 I				
				proline synthetase co-transcribed (bacterial homolog)	2.193
	133601 3		Hs.75155		2.193
65	119904 \			ESTs; Weakly similar to !! ALU SUBFAMILY SP WARNING ENTRY !! [H.sapiens]	
65	100348 [transducer of ERBB2; 2 (TOB2)	2.185
		AA351779	Hs.200334		2.18
	127793		Hs.30445	ESTs; Weakly similar to transcription regulator Staf-50 [H.sapiens]	2.178
	105149	AA169253		ESTs	2.177
		A A405648		zw39g8.s1 Soares_total_fetus_Nb2HF8_9w H sapiens cDNA clone IMAGE:772470	
					*

				•	
	111836	R36228	Hs.25119	ESTs	2.175
		R16759		ribosomal protein S5 pseudogene 1	2.175
		AA489697	Hs.145053		2.175
		F11087	Hs.239666		2.175
5		X94612		protein kinase; cGMP-dependent; type II	2.161
3		AA043223	Hs.4815	nudix (nucleoside diphosphate linked molety X)-type motif 3	2.157
		AA443828	Hs.25324	ESTs	2.157
		AA478446		KIAA1096 protein	2.156
		M19309		troponin T1; skeletal; slow	2.155
10		C06270	Hs.8078	Homo sapiens mRNA; cDNA DKFZp586L081 (from clone DKFZp586L081)	2.155
10		AA056012	Hs.9552	binder of Arl Two	2.151
		AA393755.		ESTs; Highly similar to CGI-62 protein [H.sapiens]	2.15
		U33921	113.117.211	HSU33921 Clontech adult lung cDNA library (HL1158a) Homo sapiens cDNA	2.15
		F10638	He 12/132	Homo sapiens clone 24407 mRNA sequence	2.15
15		Y10032		serum/glucocorticold regulated kinase	2.15
13		AA448710	Hs.41327		2.15
		AA399164		ESTs; Moderately similar to !! ALU SUBFAMILY SQ	2.137
				guanylate cyclase 1; soluble; alpha 3	2.137
		X66534 AA652238	Hs.199726		2.135
20				Homo sapiens BAC clone RG054D04 from 7q31	2.134
20		AA446121		KIAA0917 protein	2.132
		AA458770	Hs.82960	ESTs _	2.128
		AA121270 AA465341	Hs.99640		2.126
				adaptor-related protein complex 1; beta 1 subunit	2.125
25		L13939		adaptor-related protein complex 1; sigma 2 subunit	2,125
25		D31120	Ho 42450	Homo sapiens mRNA; cDNA DKFZp586C1817 (from clone DKFZp586C1817)	2,125
		AA039595 AA435798	Hs.98835		2.125
		H01992		KIAA1102 protein	2.125
		H17490	Hs.7905	ESTs; Highly similar to sorting nextn 9 [H.saplens]	2.123
30			He 102/12	ESTs; Modify smir to !! ALU SUBFAMILY J WARNING ENTRY !! [H.sapiens]	2.123
50		N69666 AA621202	Hs.7946	DKFZP586D1519 protein	2.12
		N71935		multiple PDZ domain protein	2.12
		U63717		osteodast stimulating factor 1	2.118
		AA405263	Hs.181400		2.109
35		H38148	Hs.32775		2.108
<i>JJ</i>		AA521186	Hs.94217		2.107
		U76189	Hs.61152		2.102
		N50073	Hs.84926		2.1
		AA419622	Hs 104800	ESTs; Weakly similar to Mouse 19.5 mRNA; complete cds [M.musculus]	2.096
40		Y09443			2.094
40		AA406293	Hs.193498		2.093
		AA398710		chloride channel 3	2.091
		F10980	Hs.184780		2.09
		N58193		ESTs: Weakly similar to 1-evidence	2.089
45		AA129931		protein phosphatase 1; catalytic subunit; gamma isoform	2.083
		N73702	Hs.238927		2.083
		R05490	Hs.7239	SEC24 (S. cerevisiae) related gene family; member B	2.079
		R40923	Hs.106604		2.078
		N47587		ESTs; Wealdy similar to TROPOMODULIN [H.sapiens]	2.076
50		Al457411	Hs.106728		2.076
•		L15309		zinc finger protein 141 (clone pHZ-44)	2.075
		F02582	Hs.14474	ESTs	2.074
		N29325	Hs.7535	ESTs; Highly similar to COBW-like placental protein [H:sapiens]	2.07
		AA478609	Hs.47278	Human Chromosome 16 BAC done CIT987SK-A-735G6	2.07
55		R16667	Hs.24752	spectrin SH3 domain binding protein 1	2.069
		T90830	Hs.15981	ESTs; Weakly similar to line-1 protein ORF2 [H.sapiens]	2.067
		D14826		cAMP responsive element modulator	2.064
		AA131866	Hs.61661	ESTs; Weakly similar to DY3.6 [C.elegans]	2.063
		R53765		KIAA0981 protein	2.063
60	125624	AA165411		zq49a01.r1 Stratagene hNT neuron (#937233) Homo sapiens cDNA clone	2.061
		N72200	Hs.13913	ESTs	2.058
		AA495830	Hs.87013		2.057
		R51361	Hs.23423		2.056
		AA424754	Hs.43149		2.056
65		AA599219		ESTs; Weakly similar to ALR [H.sapiens]	2.056
-		H30270	Hs.165062		2.054
		AA262354	Hs.186648		2.054
		N59249	Hs.48349		2.052
		AA293194	Hs.3737	ESTs	2.052

	132617	AA171913	Hs.5338	carbonic anhydrase XII	2.05
		N36167	Hs.28274	ESTs	2.05
		T64438	Hs.11449	DKFZP564O123 protein	2.05
_	122785	AA459978	Hs.99508		2.05
5	107203	D20426	Hs.5656	EST	2.05
		AA291321		ESTs; Moderately similar to KIAA1006 protein [H.sapiens]	2.046
		D82675		Homo sapiens clone 25007 mRNA sequence	2.042
		R43845	Hs.64595		2.04
10		AA600253	Hs.55601		2.04
10		AA526849 AA279442	Hs.82109	syndecan 1 protein kinase C; nu	2.039 2.037
		W70205	Hs.43670		2.037
		M21494		creatine kinase; muscle	2.036
		AA609943	Hs.32793		2.034
15		H89112		yw25e5.s1 Morton Fetal Cochlea Homo sapiens cDNA clone IMAGE:25328	2.034
	132906	AA142857	Hs.234896	ESTs; Highly similar to geminin [H.sapiens]	2.031
	105479	AA255546	Hs.23467		2.027
		U04898	Hs.2156	RAR-related orphan receptor A	2.027
20		W80363	Hs.58446		2.024
20		R46482	Hs.106875		2.024
		AA041548		KIAA0573 protein	2.023
		N50114 W37999	Hs.128704 Hs.24336		2.017 2.017
		AA281257	Hs.125868		2.014
25		AA062731		thyroid hormone receptor-associated protein; 150 kDa subunit	2.012
		AA278846	Hs.187634		2.011
	113933	W81362	Hs.30567	ESTs	2.011
	123072	AA485041	Hs.104308	ESTs	2.009
00		AA609323	Hs.112689		2.008
30		H67749	Hs.161022		2.003
		X69398		CD47 antigen (Rh-related antigen; integrin-associated signal transducer)	1.995
		Y07755 N53378	Hs.38991 Hs.22543		1.995 1.995
		AA251797	113.22040	zs11/3.s1 NCl_CGAP_GCB1 Homo sapiens cDNA clone	1.989
35		R53457	Hs.26040	ESTs; Weakly similar to fatty acid omega-hydroxylase [H.sapiens]	1.989
-		Z38652		ESTs; Weakly similar to TYL [H.sapiens]	1.988
		AA151005		sperm surface protein	1.988
	106320	AA436608		ESTs .	1.988
40		AA147224	Hs.71814		1.986
40		AA401633	Hs.22380		1.982
		AA157911	Hs.72200		1.982
		N66769 AA281886	Hs.82781 Hs.88923		1.975 1.975
		AA279060		B-cell CLL/lymphoma 10	1.974
45 .		AA719776	113.133310	zh38g04.s1 Soares_pineal_gland_N3HPG Homo sapiens cDNA clone IMAGE:4143	
	109547		Hs.26966		1.973
		AA805726	Hs.220509		1.969
	101266	L36645	Hs.73964	EphA4	1.966
~~		AA037467	Hs.30340		1.965
50		AA428240	Hs.126083	==+*	1.962
		R93696	Hs.169882	=	1.961
		AA255538	Hs.190504		1.959
		AA458578 AA251129	Hs.24416		1.956 1.953
-55	134740			oploid receptor; kappa 1	1.95
.55		AA210700		Homo sapiens mRNA; cDNA DKFZp564P056 (from clone DKFZp564P056)	1.95
		H93043	Hs.107070		1.95
		U36922		Human fork head domain protein (FKHR) mRNA, 3' end	1.948
	109441	AA228100	Hs.86998	nuclear factor of activated T-cells 5	1.946
60	127364	AA179573	Hs.90061	progesterone binding protein	1.942
		AA227498	Hs.3623	ESTs	1.942
	130672		Hs.177	phosphatidylinositol glycan; class H	1.942
		D45332	Hs.6783	ESTs	1.94
65		R62589 AA258063	Hs.167419 Hs.23438		1.939 1.937
UJ		AA490969	Hs.168147		1.936
	118873		Hs.44577		1.936
	114124			ESTs; Highly similar to KIAA0886 protein [H.sapiens]	1.934
		AA255486	Hs.88045		1.933
				•	

		H93463	Hs.124777		1.931
		AA236209	Hs.187626		1.931
		T56013		3-hydroxy-3-methylglutaryl-Coenzyme A synthase 1 (soluble)	1.929
_		AA424814	Hs.187509		1.927
5		R23241		STAT induced STAT inhibitor-2	1.925
		H06245	Hs.106801		1.925
		AA219699		KIAA0929 protein Msx2 interacting nuclear target (MINT) homolog	1.913
		H82165	Hs.40334		1.911
10		AA369027	Hs.71414		1.905
10		W44928	Hs.4878	ESTS	1.905
		AA070906	11- 404400	zm66d1.s1 Stratagene neuroepithelium (#937231) Homo sapiens cDNA clone	1.904
		AA251875		ESTs; Weakly similar to Gag-Pol polyprotein [M.musculus]	1.903
		D80063	Hs.241673		1.901
15		AA399371		ESTs; Weakly similar to zinc finger protein SALL1 [H.sapiens]	1.9
13		AA401804	Hs.114574		1.896
		F01831	Hs.14838		1.894
		W72982	Hs.58262		1.894
		AA428090	Hs.26102		1.893 1.891
20		C17938		Homo sapiens mRNA; cDNA DKFZp564O0122 (from clone DKFZp564O0122)	1.891
20		AA278907 AA461195	Hs.24549 Hs.99580		1.887
		W35390	Hs.55533		1.886
		AA134289		Homo sapiens BAC clone RG114B19 from 7g31.1	1.886
	125629	AA418069		natural killer-tumor recognition sequence	1.886
25	110611	H66947		ESTs; Highly similar to gene ERCC5 protein [H.sapiens]	1.885
		N22569	Hs.43215		1.884
		N92239			1.881
		AA013312	Hs.64988		1.881
	120285	AA182882		titin-cap (telethonin)	1.878
30	112724	R91753	Hs.17757		1.878
	103121	X63679	Hs.4147	translocating chain-associating membrane protein	1.875
	124381	N26765	Hs.109008		1.875
		N20468		ESTs; Weakly similar to putative p150 [H.sapiens]	1.875
0.5		AA279991		ESTs; Weakly similar to trithorax homologue 2 [H.sapiens]	1.875
35		N69113	Hs.110855		1.875
		AA285079	Hs.190474		1.873
		AA600012		ESTs; Moderately similar to KIAA0400 [H.sapiens]	1.872 1.872
		AA381902 AA258366		RNA binding protein ras GTPase activating protein-like	1.872
40		U26726	Hs.1376	hydroxysteroid (11-beta) dehydrogenase 2	1.87
70		M10098		ol: 18S ribosomal RNA	1.868
		AA191353		ESTs; Weakly similar to KiAA0970 protein [H.sapiens]	1.867
		S72869		DNA segment; single copy; probe pH4 (transforming sequence; thyroid-1;	1.867
		R44479		KIAA0552 gene product	1.866
45		N26175	Hs.93405		1.864
		AA053027	Hs.191797	ESTs	1.863
	123541	AA608794	Hs.112592	ESTs	1.863
	124890	R78618		ESTs; Wealty similar to RAS-RELATED PROTEIN RAB-8 [H.sapiens]	1.862
	105299	AA233511		ATP-binding cassette; sub-family G (WHITE); member 2	1.861
50	103560	Z20656	Hs.182787	myosin; heavy polypept 6; cardiac muscle; alpha (cardiomyopathy; hypertrophic 1)	
		T33637	Hs.6841	ESTs ·	1.86
		AA235040	Hs.107283		1.859
		AA243523		ESTs -	1.858
55		AA620381	Hs.70488		1.857
22		AA084323	Hs.68138		1.857 1.856
		W85812 H97678	Hs.187554 Hs.31319		1.856
		AA412087		EST; Highly smlr to prot inhibitor of activated STAT prot PIASx-alpha [H.sapiens]	1.853
		R44840	Hs.21303		1.852
60		N67317	Hs.50150		1.852
		AA004955	Hs.60015		1.851
		D87446		KIAA0257 protein	1.85
		AA287312	Hs.191648		1.85
		AA417078	Hs.193767		1.843
65		N26011		ESTs	1.843
	132084	Y12394	Hs.3886	karyopherin alpha 3 (importin alpha 4)	1.843
		N48593	Hs.121820	ESTs	1.841
		AA173440	Hs.193919		1.838
	127226	AA731036	Hs.3463	ribosomal protein \$23	1.838

	111837	R36447	Hs.24453	ESTs	1.835
		M64174	Hs.50651	Janus kinase 1 (a protein tyrosine kinase)	1.834
•		AA018937	Hs.128629		1.833
	102332	U35637		Human nebulin mRNA, partial cds	1.83
5	126579	W72979	Hs.146082	ESTs	1.83
	102341	U37122	Hs.8110	adducin 3 (gamma)	1.83
		Z39848	Hs.12079	ESTs	1.828
		D17532	Hs.316	DEAD/H (Asp-Glu-Ala-Asp/His) box polypeptide 6 (RNA helicase; 54kD)	1.823
10		AA136521	Hs.71148	ESTs; Weakly similar to putative p150 [H.sapiens]	1.823
10		AA255566	Hs.42484	Homo sapiens mRNA; cDNA DKFZp564C053 (from clone DKFZp564C053)	1.823
		AA609008	Hs.183232		1.822
	100131	D12485	Hs.11951	phosphodiesterase l/nucleotide pyrophosphatase 1 (homologous to mouse Ly-41 antigen)	1.822
	124163	H30539	Hs.189838		1.821
15		N59859	Hs.48443		1.821
		AA016021		DKFZP434K151 protein	1.82
	100357	D78156		RAS p21 protein activator 2	1.82
		AA489016	Hs.91216	ESTs; Highly similar to partial CDS; human putative tumor suppressor [H.sapiens]	1.82
20		R54112	Hs.128697		1.817
20		AA453255	Hs.6968	ESTS	1.817
		Z41589 N72253	Hs.238246	ESTs; Moderately similar to H1 chloride channel [H.sapiens]	1.815 1.813
		N30068	Hs.15347		1.812
		AA422123	Hs.42457		1.811
25		AA055404		ESTs; Weakly similar to !! ALU SUBFAMILY J WARNING ENTRY !! [H.sapiens]	1.253
	115864	AA432080	Hs.81200		1.81
•	129737	AA056140	Hs.122684	ESTs	1.81
		N53158	Hs.102682		1.809
20		HG3740-HT40		Basic Transcription Factor 2, 34 Kda Subunit	1.806
30		AA421053 AA287596	Hs.34395	ESTs zs52h09.s1 NCL_CGAP_GCB1 H sapiens cDNA clone IMAGE:701153	1.806 1.804
		AA456635	Hs.78524	ESTs	1.804
		Z39050	Hs.21963	ESTs	1.804
		N59764	Hs.5398	quanine-monophosphate synthetase	1.803
35	119135	R49548	Hs.169681	death effector domain-containing	1.802
		N91087	Hs.28728		1.801
		AA177138	Hs.161671		1.8
		N25427 Z25535	Hs.108812	nucleoporin 153kD	1.8 1.8
40		AA406367	Hs.15973	ESTs	1.8
10		H22372	Hs.163586		1.799
		AA397915 ·	Hs.77221	choline kinase	1.798
	105777	AA348412	Hs.23096	ESTs	1.797
		H19480	Hs.174309		1.796
45		AA130273	Hs.7584	ESTs; Weakly similar to hypothetical protein; similar to [H.sapiens]	1.796
		AA251330 AA279757	Hs.28248	ESTs ESTs; Weakly similar to BACN32G11.d [D.melanogaster]	1.795 1.794
		L13698	Hs.67466 Hs.65029	growth arrest-specific 1	1.794
		N48674	Hs.23796	Human DNA sequence from done 1052M9 on chromosome Xq25. Contains the	1.792
50		D13540	Hs.22868	protein tyrosine phosphatase; non-receptor type 11	1.791
		AA331157		EST35035 Embryo, 6 week, subtracted (total cDNA) I Homo sapiens cDNA	1.79
		Z38878	Hs.24979	ESTs	1.79
		Al096717		KIAA0525 protein	1.788
55		N66818	Hs.42179		1.787
55		R63925 N69682	Hs.28464	SC35-interacting protein 1	1.787 1.786
		AA600057	Hs.70266		1.784
		R40096	Hs.176578		1.784
		T89386		KIAA0606 protein; SCN Circadian Oscillatory Protein (SCOP)	1.783
60		AA262710		KIAA0627 protein	1.783
		AA489020	Hs.193424		1.782
		AA441792		chord domain-containing protein 1	1.781
		HG2463-HT25 AA374532	3 9	Guanine Nucleotide-Binding Protein G25k EST86676 HSC172 cells I Homo sapiens cDNA 5' end, mRNA sequence	1.779 1.778
65		AA436475	Hs.190104		1.777
30		AA151771			1.776
	107601	AA004636	Hs.50223		1.776
		W68255	Hs.27194	DKFZP434K171 protein	1.776
	118449	N66413	Hs.172466	ESTs; Weakly similar to KIAA0775 protein [H.sapiens]	1.776

	107060	AA034030	Un 155010	mathulmalanul Caanama A mutaan	1.775
		AA342079	Hs.252055	methylmalonyl Coenzyme A mutase	1.775
		T16305	Hs.49349		1.775
		AA406105	Hs.5344	beta-site APP-cleaving enzyme	1.774
5		AA373091	Hs.93832	adaptor-related protein complex 1; gamma 1 subunit Homo sapiens clone 24483 unknown mRNA; parital cds	1.774
5		AA428379	Hs.24870		1.773
			Hs.172635		1.773
		N26777 AA435664	Hs.8583		1.773
		AA436705	Hs.28020	similar to APOBEC1	1.772
10		N93797	Hs.3090		1.772
10		AA479166	Hs.105633	EphB1	1.772
		AA291946	Hs.42736		1.771
		AA180208	Hs.16606	ESTs; Highly similar to CGI-32 protein [H.sapiens]	1.767
		AA188618		ariadne; Drosophila; homolog of	1.766
15		AA398290	Hs.21965	ESTs	1.764
13		M86917	Hs.24734	oxysterol binding protein	1.764
		D13628	Hs.2463		1.764
		R07335	F15.2403	angiopoietin 1 ye96c1.s1 Soares fetal liver spleen 1NFLS Homo sapiens cDNA clone	1.763
		AA442257	Hs.192076		1.762
20		H02566		Homo sapiens mRNA; cDNA DKFZp434N174 (from clone DKFZp434N174)	1.761
20		AA428069		KIAA1046 protein	1.758
		AA620782	Hs.23247		1.757
		AA338960	Hs.28170		1.756
		AA435536	Hs.24336		1.756
25		AA304566	Hs.3542	ESTs	1.756
23		AA234945	Hs.11360		1.756
		N50112	Hs.47023		1.754
		AA599472		succinate-CoA ligase; GDP-forming; beta subunit	1.754
		R45963		ESTs; Weakly similar to ORF2 [M.musculus]	1.753
30		D42047		KIAA0089 protein	1.753
		N47938		yy84a09.s1 Soares_multiple_sclerosis_2NbHMSP Homo sapiens cDNA clone	1.751
		AF006082	Hs.62461	ARP2 (actin-related protein 2; yeast) homolog	1.751
		AA455917	Hs.50785	SEC22; vesicle trafficking protein (S. cerevisiae)-like 1	1.75
	110759	N21671	Hs.19025		1.75
35		AA424515	Hs.33264		1.75
		U43701		ribosomal protein L23a	1.75
	115867	AA432162	Hs.165986	DKFZP586B2022 protein	1.749
		AA194075	Hs.99908	nuclear receptor coactivator 4	1.747
	113783	W19222	Hs.7041	ESTs; Weakly similar to !! ALU SUBFAMILY SQ WARNING ENTRY !! [H.sapiens]	1.747
40	134898	X98330	Hs.90821		1.745
•	132215	T10132	Hs.4236	KIAA0478 gene product	1.744
		AB002346	Hs.61289	synaptojanin 2	1.743
		AA461556	Hs.202949	KIAA1102 protein	1.743
4.5		AA284252	Hs.58372	ESTs	1.743
45		AA236545	Hs.54973		1.742
		AA470941	Hs.143162		1.741
		Z38807	Hs.22870		1.739
		T88908	Hs.189746		1.738
50		F10577	Hs.70312		1.735
50		R51476	Hs.194524	yg76f04.r1 Soares infant brain 1NIB Homo sapiens cDNA clone	1.733
		AA279654		20.0	1.733
		U18242		calcium modulating ligand	1.732
		H88157	Hs.41105 Hs.23361		1.731 1.731
55		R22212 H72240		ESTs; Moderately similar to KIAA0745 protein [H.sapiens]	1.731
<i>JJ</i>		AA412063	Hs.6065	ESTs	1.728
		AA101984		G-protein coupled receptor	1.726
		R12581	Hs.191146		1.726
		L76703		protein phosphatase 2; regulatory subunit B (B56); epsilon isoform	1.725
60		AA489086	Hs.36545		1.725
00		N52136	Hs.93828		1.725
		AA400422	Hs.55896		1.725
		AA404995	Hs.192480		1.725
		R27296	Hs.23240		1.725
65		AA479181	Hs.186726		1.725
		U44754		small nuclear RNA activating complex; polypeptide 1; 43kD	1.724
		H29050	Hs.24096		1.722
		AA127395	Hs.222414		1.722
		H88477	Hs.191178		1.721
					-

	120271	AA176404	Hs.111092	ESTs; Weakly similar to ZINC FINGER PROTEIN 136 [H.sapiens]	1.72
		D28915		Interferon-induced; hepatitis C-associated microtubular aggregate prot (44kD)	1.719
		W69459		sex comb on midleg (Drosophila)-like 1	1.719
			Hs.8750	ESTs	
5		W73367		==	1.717
3		AA055475		clathrin; light polypeptide (Lca)	1.717
		AA281290		ESTs; Weakly similar to BC331191_1 [H.saplens]	1.717
	134891	F03517	Hs.90787	ESTs	1.716
	106219	AA428567	Hs.26613	Homo saplens mRNA; cDNA DKFZp586F1323 (from clone DKFZp586F1323)	1.715
	116372	AA521311	Hs.13854	ESTs	1.713
10	107570	AA001870	Hs.237323	N-acetylglucosamine-phosphate mutase; DKFZP434B187 protein	1.713
		AA427816	Hs.11803		1.712
		W31479	Hs.129051		1.712
					1.712
		AA085676	Hs.6763	KIAA0942 protein	
15		J04813		cytochrome P450; subfamily IIIA (niphedipine oxidase); polypeptide 5	1.711
15		D20899		Homo sapiens mRNA; cDNA DKFZp564G022 (from clone DKFZp564G022)	1.711
		AA766511	Hs.128848		1.71
	116089	AA455933	Hs.41324		1.709
	123337	AA504153	Hs.132797	ESTs; Weakly similar to ORF YGL050w [S.cerevisiae]	1.708
	123619	AA609200	Hs.162686	ESTs	1.708
20	104781	AA026617	Hs.21610	ESTs; Highly similar to BAI1-associated protein 1 [H.saplens]	1.707
		AA256468	Hs.88148		1.705
		N49408		KIAA0853 protein	1.705
		T57570	Hs.77039		1.704
	4	N91273	Hs.27179	• The state of the	1.702
25					1.702
23		L36644	Hs.31092		
		F08925	Hs.48610		1.7
		N67192	Hs.49476		1.7
		F02488	Hs.21917		1.7
	128499	AA487503	Hs.100636		1.698
30	120780	AA342337	Hs.241569	ESTs; Modtly smir to !! ALU SUBFAMILY SQ WARNING ENTRY II [H.sapiens]	1.697
	132920	L06133	Hs.606	ATPase; Cu++ transporting; alpha polypeptide (Menkes syndrome)	1.696
	135037	U77948	Hs.184122	general transcription factor II; i	1.696
	110024	H11297	Hs.31050	ESTs	1.695
		AA329274	Hs.82911	protein tyrosine phosphatase type IVA; member 2	1.694
35		U24685		Human anti-B cell autoantibody IgM heavy chain variable V-D-J region (VH4)	
55	IULLEU		113.140220	gene; done E11; VH4-63 non-productive rearrangement	1.694
	100710	AA205862	Hs.7942	ESTs	1.694
					1.692
		M27492	Hs.82112	interleukin 1 receptor, type I	
40		AA435551	Hs.30824	ESTs	1.691
40	116826	H58691	Hs.8215	ESTs; Weakly similar to double-stranded RNA-binding nuclear	
				protein DRSBP76 [H.sapiens]	1.69
	135339	D59269	Hs.127842	Homo sapiens mRNA full length insert cDNA clone EUROIMAGE 783648	1.69
	118250	N62602		yz75b6.s1 Soares_multiple_sclerosis_2NbHMSP Homo sapiens cDNA done	
				IMAGE:288851 3' similar to contains Alu repetitive element;, mRNA sequence	1.689
45	106470	AA450116	Hs.186180	ESTs	1.688
	108203	AA057678	Hs.63408	ESTs	1.687
		W70313	Hs.126906	ESTs	1.686
		D51228	Hs.79404		1.683
		AA481392	Hs.105166		1.683
50		AA011616	Hs.184086		1.681
20		*****			1.678
		M28209	HS.2307 10	RAB1; member RAS oncogene family	
	-	U76638		BRCA1 associated RING domain 1	1.677
		AA256386		• • • • • • • • • • • • • • • • • • • •	1.676
		N67277	Hs.9403	ESTs	1.676
55	105937	AA404342	Hs.173531	ESTs	1.675
	114118	Z38520	Hs.175930	ESTs	1.675
	109203	AA190634	Hs.108787	endoplasmic reticulum membrane protein	1.675
	125245	W86608	Hs.7243	ubiquitin specific protease 24	1.675
		X06956		tubulin; alpha 1 (testis specific)	1.675
60		AA262925		cleavage stimulation factor, 3° pre-RNA; subunit 3; 77kD	1.674
		U63289		CUG triplet repeat; RNA-binding protein 1	1.674
		F10108	Hs.183333		1.673
					1.673
		D63876		KIAA0154 protein	
65		AA402937	Hs.103238		1.671
65		AA001386	Hs.59844	COIS	1.671
		AA243139	Hs.4863	Homo sapiens done 25088 mRNA sequence	1.669
		N58172	Hs.109370		1.668
		H92575		ESTs; Weakly similar to II ALU SUBFAMILY SQ WARNING ENTRY II [H.sapiens]	1.668
	126809	AA743475	Hs.171693	ESTs	1.667

	106095	AA419547	Hs.11713		1.66
	101754	M77142	Hs.239489	3 TIA1 cytotoxic granule-associated RNA-binding protein	1.66
	105188	AA192306	Hs.23926	ESTs	1.66
_	113582	T91371	Hs.16824		1.66
5	119559	W38197		Accession not listed in Genbank	1.66
	119961	W87535	Hs.59015	ring finger protein 9	1.65
	123255	AA490890	Hs.105273		1.65
	111078	N59230	Hs.186574	ESTs .	1.65
	113082	T40528	Hs.8246	ESTs	1.65
10	119589	W44692	Hs.124177	' ESTs	1.65
	104308	D53639		ribosomal protein \$26	1.65
	103073	X59417	Hs.74077	proteasome (prosome; macropain) subunit; alpha type; 6	1.65
	124424	N35314	Hs.107265	S ESTs	1.65
	128890	AA096157	Hs.182364	ESTs; Weakly similar to 25 kDa trypsin inhibitor [H.sapiens]	1.65
15	119400	T92767		ye27d06.s1 Stratagene lung (#937210) Homo sapiens cDNA clone	
				IMAGE:118955 3', mRNA sequence.	1.65
		AA486868	Hs.29802		1.65
		N62339	Hs.180532	heat shock 90kD protein 1; alpha	1.649
00		N67954	Hs.49413		1.648
20		AA476307	Hs.194035	KIAA0737 gene product	1.647
		X60708	Hs.44926	dipeptidylpeptidase IV (CD26; adenosine deaminase complexing protein 2)	1.647
		U69140		fasciculation and elongation protein zeta 2 (zygin II)	1.646
		T15530	Hs.221439	•	1.646
25		AA056263	Hs.132747		1.64
25		AA579377	Hs.180532	heat shock 90kD protein 1; alpha	1.644
		AA007595	Hs.220937		1.642
		N79820	Hs.50854		1.64
		D85423		Homo sapiens mRNA for Cdc5, partial cds	1.64
20		AA284865		KIAA1040 protein	1.639
30		AA460128	Hs.5074	similar to S. pombe dim1+	1.639
		AA034002	Hs.76359		1.639
		AA447083	Hs.134522		1.637
	110370	AA521256	MS.236204	ESTs; Moderately similar to NUCLEAR PORE COMPLEX	
35	100044	E04477	Ha 204000	PROTEIN NUP107 [R.norvegicus]	1.631
55	109644	F04477	ns.204802	ESTs; Moderately similar to GLYCERALDEHYDE 3-PHOSPHATE	4
	103427	V07202		DEHYDROGENASE; LIVER [H.sapiens]	1.627
	132186		He 991040	H.sapiens mRNA for Ptg-12 protein KIAA1038 protein	1.627
	131428		He 26710	PR domain containing 2; with ZNF domain	1.626
40		AA649257	Hs.188602		1.626
		AA039568	Hs.188083	·	1.625 1.625
		AA400857	Hs.97509		1.625
		AA446885		ESTs; Moderately similar to ZINC FINGER PROTEIN 141 [H.sapiens]	1.625
	110632		Hs.171635		1.624
45	111389			ESTs; Weakly similar to L82A [D.melanogaster]	1.624
	112449		Hs.124186	ring finger protein 2	1.623
	113070		Hs.6298	ESTs	1.622
	107229		Hs.34644		1.618
	132710			protease Inhibitor 5 (maspin)	1.617
50	124664	N94814	Hs.33540	ESTs; Weakly similar to KIAA0765 protein [H.sapiens]	1.617
	130166	AA350690	Hs.151411	KIAA0916 protein	1.616
•	125040	T78451	Hs.199961	ESTs	1.615
	132972	H39627		ESTs; Weakly similar to !! ALU SUBFAMILY SB WARNING ENTRY !! [H.sapiens]	1.615
	115873	AA433916		heat shock 70kD protein 4	1.611
55	120408	AA235045	Hs.190151	ESTs	1.61
	120934	AA383773	Hs.191500	ESTs	1.61
	115259	AA279071	Hs.13453	splicing factor 3b; subunit 1; 155kD	1.609
	134330	D20113	Hs.8185	ESTs; Highly similar to CGI-44 protein [H.sapiens]	1.607
	115117	AA256492	Hs.49007	poly(A) polymerase	1.606
50	125162	W44682	Hs.109896		1.605
	103946	AA285246	Hs.111650	ESTs; Weakly similar to Prt1 homolog [H.sapiens]	1.604
	133389	AA166917	Hs.72639	ESTs .	1.603
	115528	AA342301	Hs.53929	ESTs; Weakly similar to !! ALU CLASS B WARNING ENTRY !! [H.sapiens]	1.602
	129704	W81301	Hs.12064	ubiquitin specific protease 22	1.602
55	109313	AA206800		ESTs; Moderately similar to zinc finger protein dp [H.sapiens]	1.601
	130457		Hs.155976		1.6
		AA485211	Hs.190046		1.6
	115113	AA256460	Hs.44610	ESTs	1.6
	117731	N46433	Hs.46609	ESTs	1.6

	123344	AA504338	Hs.171857		1.599
	131798	X86098	Hs.3238	adenovirus 5 E1A binding protein	1.597
	125370	AA256743		KIAA0092 gene product	1.596
	114918	AA236813	Hs.72324	ESTs; Highty similar to unknown [H.sapiens]	1.596
5		AA160805	Hs.199832	ESTs	1.596
	105103	AA151593	Hs.10130		1.594
	125004	T60120		yb68f02.s1 Stratagene ovary (#937217) Homo sapiens cDNA clone	
				IMAGE:76347 3', mRNA sequence.	1.592
	105658	AA282914	Hs.10176	ESTs	1.589
10	110455	H52172		yt85e8.s1 Soares_pineal_gland_N3HPG Homo sapiens cDNA clone	
				IMAGE:23111 3' similar to contains Alu repetitive element, mRNA sequence	1.589
	119780	W72967	Hs.191381	ESTs; Weakly similar to hypothetical protein [H.sapiens]	1.587
	126983	AA211537		zn55d01.r1 Stratagene muscle 937209 Homo sapiens cDNA clone	
				IMAGE:562081 5', mRNA sequence.	1.586
15		AA250745	Hs.87773	protein kinase; cAMP-dependent; catalytic; beta	1.584
		AA252033	Hs.15036		1.584
		Z40251	Hs.56974		1.584
		AA428137	Hs.86434		1.581
00		AA456311	Hs.93961	ESTs; Weakly similar to !! ALU CLASS A WARNING ENTRY !! [H.sapiens]	1.581
20		AA479295		Kelch motif containing protein	1.581
		W67569	Hs.44143	ESTs; Weakly similar to SNF2alpha protein [H.sapiens]	1.58
		D80948	Hs.34922		1.58 1.58
		AA424558	Hs.9302	phosducin-like	1.579
25		AA279422	Hs.5724	ESTs	1.577
25		R27598		KIAA0797 protein	1.575
		R98173	Hs.23763		1.575
		N21680	Hs.43047		1.575
		M33772		troponin C2; fast v-myc avian myelocytomatosis viral oncogene homolog	1.575
30		AA459703 W90625	Hs.79070 Hs.58432	ESTs	1.575
30		N32157	Hs.82207		1.574
		AA452865		UDP-Gat:betaGlcNAc beta 1;4- galactosyltransferase; polypeptide 2	1.573
		AA609204	Hs.27973		1.573
		AA810215	Hs.189079		1.571
35		W72798		ESTs; Wkty smlr to cDNA EST EMBL:D32579 comes from this gene [C.elegans]	1.571
J J		AA135638	Hs.223756		1.571
		AA456112	Hs.99410		1.57
		H12636		ESTs; Weakly similar to reverse transcriptase [H.sapiens]	1.568
		AA609828	Hs.21015	ESTs: Highly similar to tetracycline transporter-like protein [M.musculus]	1.568
40		Z41366		KIAA0872 protein	1.567
		N53076	Hs.5996	ESTs	1.567
		AA913491	Hs.189143	ESTs; Modrtly smlr to !! ALU SUBFAMILY J WARNING ENTRY !! [H.sapiens]	1.567
	112046	R43365	Hs.22273	ESTs	1.566
	132160	AA281770	Hs.184081	seven in absentia (Drosophila) homolog 1	1.566
45	111568	R10153	Hs.20561	ESTs	1.566
	127775	H04106	Hs.179902	ESTs; Weakly similar to NG22 [H.sapiens]	1.566
	115359	AA281936	Hs.88914	ESTs	1.566
		AA425734	Hs.165066	ESTs; Weakly similar to hypothetical protein [H.sapiens]	1.565
~~	127854	AA769520		ESTs; Weakly similar to REGULATOR OF MITOTIC SPINDLE	4 000
50				ASSEMBLY 1 [H.saplens]	1.564
		AA187679	Hs.111114		1.563
		AA243012	Hs.75928	ESTs	1.562
		AA031700	Hs.251962		1.562
<i></i>		U97188	Hs.79440	IGF-II mRNA-binding protein 3	1.561
55		H95094	Hs.75187	translocase of outer mitochondrial membrane 20 (yeast) homolog	1.561
		AA281244	Hs.65300	ESTs	1.559
		T97931	Hs.18190		1.558
		AA236177	Hs.76591	KIAA0887 protein	1.558
60		T62571		microtubule-associated protein 7	1.558 1.558
60		AA039616	Hs.61933	ESTS	1.557
	104054	AA156499	Hs.8454	protein kinase; cAMP-dependent; regulatory; type II; alpha	1.557
	134351	R82074	Hs.82109 Hs.70333	syndecan 1 ESTs	1.553
	100000	D51401 AA490899	Hs.24462	EQTe	1.553
65					1.552
UJ		N74075 W20016	Hs.94293	ESTs; Weakly similar to ZINC FINGER PROTEIN 83 [H.sapiens]	1.55
		AA438720	Hs.65487		1.55
	107260	AA025234	Hs.61260		1.55
		N21407	Hs.257325	FSTs	1.55
	10-10-0	146 1701	لنتان البحرب ا		

	109703	F09684	Hs.24792		1.55
	120288	AA187938	Hs.55189	ESTs; Weakly similar to F25B5.3 [C.elegans]	1.548
	106356	AA443277	Hs.31034		1.548
		AA235627	Hs.11171	APG5 (autophagy 5; S. cerevislae)-like	1.547
5		D11961	Hs.77823	ESTs	
3					1.546
		Al400862	Hs.142607		1.546
		Z38909	Hs.22265		1.545
		M86546		pre-B-cell leukemia transcription factor 1	1.544
		T93630		ESTs	1.542
10	104896	AA054228	Hs.23165	ESTs	1.541
	114477	AA032013	Hs.144260	EST	1.54
	110731	H98653	Hs.188006	KIAA0878 protein	1.54
		Z38501	Hs.8768	ESTs; Wkly smlr to !! ALU SUBFAMILY SQ WARNING ENTRY !! [H.sapiens]	1.538
		L07044		Homo sapiens calcium/calmodulin-dependent protein kinase II mRNA; partial cds	1.538
15		W60186	He 160/87	Kreisler (mouse) maf-related leucine zipper homolog	1.537
13		W24957	Hs.16281		1.557
	130303	VV24801	MS. 10201	ESTs; Moderately similar to similar to C.elegans protein	4 503
				encoded in cosmid T20D3 [H.sapiens]	1.537
		AA088851	Hs.75744	S-adenosylmethionine decarboxylase 1	1.537
•	106450	AA449469	Hs.11859	ESTs	1.536
20	104120	AA429838	Hs.89519	KIAA1046 protein	1.536
	100533	HG1879-HT19	19	Ras-Like Protein Tc10	1.535
	130664	R09049	Hs.17625	ESTs	1.535
	127122	AA279153	Hs.190049		1.535
		T03391	Hs.8087	ESTs	1.535
25		AA418662	Hs.44625	ESTs	1.535
23					
		AA286941	Hs.43691	ESTs	1.533
		T59442	Hs.100445		1.532
		U30888	Hs.75981	ubiquitin specific protease 14 (tRNA-guanine transglycosylase)	1.532
20		AA426299	Hs.98510	ESTs T	1.532
30		Z41747	Hs.469	succinate dehydrogenase complex; subunit A; flavoprotein (Fp)	1.531
	132944	AA054515	Hs.6127	ESTs; Weakly similar to prostate-specific transglutaminase [H.sapiens]	1.53
	111199	N68210	Hs.29822	ESTs	1.53
	113494	T88878	Hs.258738	ESTs	1.529
	129515	AA490882	Hs.112227		1.528
35		AA156049	Hs.65490		1.528
-		AA027163	Hs.7942	ESTs	1.526
		AA279408	Hs.25866		
					1.526
		U67156		mitogen-activated protein kinase kinase kinase 5	1.526
40		Z40758		DKFZP434K151 protein	1.525
40		T03488	Hs.4842	ESTs	1.525
		AA525014	Hs.162115		1.525
			Hs.129887	cadherin 19 (NOTE: redefinition of symbol)	1.525
	119859	W80702		ESTs .	1.525
	129944	L00389	Hs.1361	cytochrome P450; subfamily I (aromatic compound-inducible); polypeptide 2	1.524
45	118864	N89670		ESTs; Weakly similar to Su(P) [D.melanogaster]	1.523
	123964		Hs.210115		1.523
			Hs.166459		1.522
			Hs.134173		1.522
		X17059			1.521
50			Hs.12396	N-acetyltransferase 1 (arylamine N-acetyltransferase)	
50		.W58461	HS. 12390		1.521
	12/093	AA768241		oa72d02.s1 NCI_CGAP_GCB1 Homo sapiens cDNA clone	
				IMAGE:1317795 3', mRNA sequence.	1.521
			Hs.4113	S-adenosylhomocysteine hydrolase-like 1	1.521
	125303	Z39821	Hs.107295	ESTs	1.52
55	132697	AA281951	Hs.5518	Homo sapiens mRNA; cDNA DKFZp566J2146 (from clone DKFZp566J2146)	1.52
	117086	H93135	Hs.41840	ESTs	1.519
	113355	T79203	Hs.14480	ESTs	1.518
				ESTs	1.518
				EST	1.518
60				RAB28; member RAS oncogene family	1.517
50				myosin X	
	132968				1.515
	117035			ESTs	1.515
	116781		Hs.52132		1.513
~~			Hs.118531		1.513
65	130214	H78003	Hs.15266	ESTs	1.513
	134700	AA481414	Hs.8868	golgi SNAP receptor complex member 1	1.512
				ESTs	1.508
	126257			tumor necrosis factor receptor superfamily; member 10b	1.508
		AA806808	Hs.118797	ubiquitin-conjugating enzyme E2D 3 (homologous to yeast UBC4/5)	1.508

	113837	W57698	Hs.8888	ESTs	1.507
	114317	Z41038	Hs.469	succinate dehydrogenase complex; subunit A; flavoprotein (Fp)	1.507
	100311	D50640	Hs.184653	phosphodiesterase 3B; cGMP-inhibited	1.507
_	126802	AA947601	Hs.97056	ESTs	1.508
5	128661	R82837	Hs.103329	KIAA0970 protein	1.508
	134194	AA233231	Hs.79828	ESTs	1.506
	108953	AA149652	Hs.42128	ESTs	1.504
	133240	D31161	Hs.68613	ESTs	1.502
	132671	X76302	Hs.54649	putative nucleic acid binding protein RY-1	1.501
10	132609	Z48923	Hs.53250	bone morphogenetic protein receptor; type II (serine/threonine kinase)	1.501
	105574	AA278678	Hs.258567	ESTs	1.5
	113718	T97782	Hs.256268	ESTs	1.5
	127824	AI208365	Hs.127811	ESTs	1.5
	130132	U55936	Hs.184376	synaptosomal-associated protein; 23kD	1.5
15	127394	AA453224		ESTs; Weakly similar to !! ALU SUBFAMILY J WARNING ENTRY !! [H.sapiens]	1.5
	100485	HG1111-HT1111		Ras-Like Protein Tc21	1.5
	101078	L04510	Hs.792	ADP-ribosylation factor domain protein 1; 64kD	1.5
	128611	AA456845	Hs.102471	KIAA0680 gene product	1.5

TABLE 12A shows the accession numbers for those primekeys lacking unigeneID's for Table 12. For each probeset we have listed the gene cluster number from which the oligonucleotides were designed. Gene clusters were compiled using sequences derived from Genbank ESTs and mRNAs. These sequences were clustered based on sequence similarity using Clustering and Alignment Tools (DoubleTwist, Oakland California). The Genbank accession numbers for sequences comprising each cluster are listed in the "Accession" column.

5

10	Pkey: CAT nui Accessio		Unique Eos probeset identifier number Gene cluster number Genbank accession numbers
15	Pkey	CAT number	Accession
	PREY	CAT HUMBER	ACCESSION
20	117040	119811_1 46956_1 18457_1	AA084524 AA339253 AW966289 AW970600 AA503323 H89218 AF086031 H89112 AA355435 NM_001516 Z30093 T28405 AW949486 AA461142 AA410532 Al652073 AA521208 Al970141 Al968234 Al026102 AA713583 AW135876 AA936614 AA770300 Al242635 AA377033 AW960263 AW607683 Al273803 AA410287 Al040513
25		3022_1	AA460838 AI803916 AW294095 AW449680 AW798677 AW675048 BE542116 AL120521 L34840 NM_003241 U31905 AI546931 AI791616 AI973065 AI792321 AI546937 AI685880 AI732835 AI682360 AA420653 AA564047 AI682323 AI824614 AI659889 AI680052 AI970887 AI623108 AA420692 AI418074 AA631018 AI810595 AW291463 AW449930 AI668908 AI970818
	100824	5_36	Al393237 Al521317 Al761348 AF025841 D43968 AW994987 L34598 AF025841 D89789 D89788 D89780 AW998932 Al971742 Al310238 X90976 AW139668 AW674280 Al365552 AA877452 AV657554 C75229 AA376077 Al798056 AW609213 W25586 H30149 BE075089 BE075190 AW580858 H99598 AA425238 AA133916 AW363478 BE158121 BE158127 AW467960 BE158135 BE158126 BE158145 N92860 AA847246 Al961688 Al361423 AA878154 AA043767 Al863712
30			AI559226 AW339007 AI371266 AI368901 AA046624 AA134739 AW449154 AA130232 AI458720 AA962511 AI700627 R70437 AW004008 AA045229 AI671572 H99599 AA043768 AI685454 AI871685 N29937 X90977 AA524240 AI142114 AI825750 AI567805 AI631365 AI347893 AA134740 F20669 AA046707 AW793216 AW963298 AW959380 AA363265 AI784593 AI268201 R69451 AV657618 AI695588
35		264197_1 27608_1 553_1	BE312163 AJ230798 AA374482 AI926059 AA622653 AI860704 BE139185 AW296884 T60238 T60120 U33921 AI190489 AA573311 AI814663 AA806761 AA765241 AA019317 AA092255 AA035405 T85079 AA890151 AI373959 T85080 BE153728 AA740848 BE080682 AL048137 AW182316 AI699468 AW274481 AW407538 AA306562 AW950024 AW949943 AL045703 AW843196 W25132 BE612794 AA304266 AW958054 H25673 AV646563 AV646573 BE172990 AW593488 AA385181 AA164998
40			Al246476 AA345406 Al277554 AA134749 AA856624 BE613247 AA299003 AL048138 AA028121 T92510 Al923835 AW020440 Al401594 Al889401 N93290 AA044247 AA028100 Al582845 AA811151 Al741811 Al925878 AA448277 AA172221 Al214783 BE220793 AA022746 Al082882 AA022849 Al928385 AA573472 Al420686 AW072902 Al799493 Al873506 Al468977 Al192079 Al468976 AA044272 AW015701 AW316979 AA933042 AA609017 Al318393 Al424571 Al934945 AA172023 AW050917 AA846180 AA134748 AL003947 Al766769 AV0A6747 AAA54585317 AW575680 Al474214 AAA64678 AM576769 AA167474 AA564678 AA56478 AA564
45			U36922 AA927064 AA868000 D62654 T91745 AW500202 AA194764 AA746346 AA130464 AW117498 AA054526 N26432 H02534 H04964 AW303367 BE300931 Al218049 Al208073 AW182749 AA983630 A147585 AA194765 AA054534 AA922720 Al436585 Al346535 AA134269 AA26923 AA897422 AA019559 AW274010 AA035406 AA917879 H99327 W32908 Al216046 AW496823 AA019414 H82288 W35284 Al936621 Al767113 AA866177 AW367874 H82398 AF032885 AW300151 AW467069 AA809346 Al188507 Al494178 AA872752 Al631631 U02310 NM_002015 AA815006 Al382453 AW197658 Al761654 Al804396 Al382221 Al813640 Al439635 Al523901 AW517242 Al221705 AW298104 AW204560 AW573095 AW028783
50	124704	292319_1	AW014650 Al766744 Al808294 Al698758 Al041809 Al766667 Al479103 AA872797 AA769305 AA765080 AA334166 Al472322 R07335 R07640
55	116988 124825 110455	185904_1 330773_1 46874_1 182217_1	AW953679 AW953680 AA244436 H82527 AA361046 AA244483 H82526 AA501669 R52088 H52576 AF085971 H52172 N99638 AW973750 AA328271 H90994 AA558020 AA234435 N59599 R94815
60	104038	154135_1 264235_1 43892_1	AW968363 AA465492 R34539 AA165411 AA374532 AA421255 BE514383 AA071273 AW247987 AW673286 BE312102 AW749824 BE071985 AW577383 BE071945 BE072005 AW577355 BE071965 AW239231 BE072000 BE071960 AW577360 AW749830 AW373020 X97303 AW999522 BE000192 BE562219
		113242_1 47721_1	BE266655 BE264970 AA074713 AA447006 AW977549 AA256038 AL365415 AW500455 AA768241 AW968097 Z17849 AA256104

	125873	10492_1	AW271838 AL133605 C01646 H29959 AA999896 D60676 AW999454 AW961176 AA315244 H14437 AW386118 N46512 AW272021 AI768516 BE466421 AI082809 AI804454 AA905101 AW173368 N38942 AW614169 AI080483 N29489 AI500550
5	125954	4457_1	AA994475 AA614464 AA707368 AA593145 AA569473 AW627815 AI828244 N63226 N42300 NM_016353 AB023584 W44753 R09585 AA382865 R23772 AI814257 AA974046 AK001608 AI935638 AW440609 AI420022 AA777386 AA806969 AI554876 AI584006 AI688556 AI688634 AI697997 AI014540 AI806683 AI741202 AW263154 AW297238 AI149951 AI589076 AW082158 AW614265 AA931887 AA781969 R09490 AA484643 AI207121 AI088390 AI538065 AI619547 AI741925 AIV2846 H40846 R93943 AW747979 AA461348 U30163 AA326023 AI535992 AW242870 AI244025 AI222558 W38425 AW473630 AI624599 AI921226 AI683152 AI096458 AI123822 AW170802 C16447 AI337674
10		1589048_1 15307_6	D25726 AW339366 AW771259 AA461174 H48372 W01626 AA305278 AA223833 110924 6443_1 AW058463 AF195766 AA680145 T86901 W60373 W60281 NM_007222 AF106862 AI000795 AA167188
15			AW884503 AW891313 AW891332 AW891312 AI984924 AI123518 N75170 AA131614 H25330 AI913358 AI742277 W25576 R58771 AW445159 AW888628 AW888627 AW274674 AI088482 N52314 N34282 AW001769 AI338943 T66784 AI288963
			AW468676 AW237528 H25289 N71690 AA610128 AI143458 AI082599 N49144 AA854773 AW663411 AW610151 N47938 AW601626 AA167189 AA918304 AA805205 BE069496 AA652836 BE069499 AI699298 AW249926 AW888578 BE567635 T10726 AW604715 D54245 D53062 D55610 D55555 AA301376 AI133498 N77788 AI936320 AW090734 AI269977 N50828 AA550814 AI421993 AI005384 N50813 D60292 D59349 AA131710 D81698 D81699
20		232161_1 29440_1	AA331156 AA331157 AA331155 U76456 NM_003256 AF057532 AA193414 AW293304 AW963378 AA313095 AI359841 AI969312 AI080163 AW448926
20			Al671136 BE466399 Al637967 Al671873 AW196583 AW071635 Al634427 AW296872 AW292470 AA193650
	127394 126879	304844_1	BE161832 AA453224 AA485772
		171841_1	D90391 M55575 Al652268 AA719776 AA524886 AW971347 AA211537
25		188975_1	AW971327 AA524988 AW628653 AA251797
		443883_1	AW976796 AA769520
		280429_1	AA432071 AA405648 AW000908 T16347
	106320	0435_1	AB028957 AL 120001 Al267678 H10928 R19844 AW970334 AA393182 F05472 F11711 H09908 N50250 Al815411 BE463679 D61468 AW970253 D60889 C15548 D61011 D60867 Al815795 AA534831 D81386 AW235039 Al382158 D81174 AA416899
30		•	AA852310 H09789 H10929 H09813 F09369 R44721 D51515 Z38456 R14004 T66255 F12148 F12139 AW351702 M85350
			Al018713 AW972450 AW972645 AA514964 T66172 F09785 F09776 AA436608 T05327 T07118 AA339352
		201515_1 11075_1	AW301608 N46706 AA649093 AA287595 AW811753 AA287596 N39260 NM_001874 J04970 T91426 AW205201 T84979 AA255727 AA847837 R02164 T91339 AV651884 AV651835 AV651350
	101020	110/5_1	AV650118 AV651338 Al272002 Al367796 AA830651 AA262112 AW151198
35	100401	24827_1	AU076696 AA219720 AL135197 AA305877 N56376 AA318063 AA130725 AW954903 BE541230 AW383312 U86753 D85423
			Al679458 Al122932 AB007892 Al583919 BE160134 F08104 R34903 F13440 AA095444 AA262453 AA191036 R17895
			T81266 BE149776 Al279537 Al143113 AA361072 AW959030 AW268817 AA811533 BE275179 Al221677 T65147 R49293 AA249176 BE000290 AA768053 F09494 BE092645 BE172099 Z41177 AA044750 Al909768 BE140795 BE140574 AW845210
			AW752452 BE243244 AA843664 Al300080 BE169032 AW189979 BE004869 AA621872 Al951772 Al678897 Al926598
40			N62813 Al350912 AW608791 Al309602 Al983138 AW875592 Al655073 AW875626 AA130606 Al370827 C75528 C75554
			AW263335 Al344426 BE004788 AA576220 AA604824 Al431405 AA749378 R38882 AW955075 AA173821 C75657
		•	AA219672 AW768408 R43141 Al431414 AA483343 Al673792 T17294 AW770187 N74285 Al476404 Al088288 AA654152 AW974864 BE617311 BE243328 BE168049
	130542	28089_3	U64675 AW167507 AW167508 BE218568 AA779360 W85722 AL044843 BE159404 AF012086 AW898611 AW898610
45		_	BE159405 BE092191 AW890826 AW369841 AW368064 AW606702 AL044731 R82691 AA419346 AA416558 H96045 AL040450 Al640531 Al808434 AL046613 AW855784 AW362469 AL048881 AL049015 AA094272 AA888908 AA417294 AW237786 R59793 AL044916 D82402 Al216854 Al079342 H96406 AL037845 Al915900 AA972133 Al478783 T31074 Z21135 Z21396 AA352182 R13918 AA430178 C17811 Al371824 Al742256 AA926801 N79156 AA350610 AA081971 N83639
50			R35544 AA312292 AW952080 N42322 AA171957 AA565297 R89207 AA504106 Al630782 AA826482 Al301579 T36241
50			AW966618 Z28426 AL043480 Al124636 AA393449 T19504 AW887823 Al289814 N53979 AL043571 Al632764 Al859613
			Al986308 Al683212 Al984499 Al133258 C05898 AW512761 Al041260 BE466240 Z19161 Al351190 N67549 Al373374 AA400873 AW440914 AW514879 AA770146 Al358754 R51113 Al283773 AA649886 T30543 D54358 R37750 T03358
			T15451 T15880 AA999689 N67396 Al056289 T65597 N62441 R89099 R00035 T85596 R61335 R00128 N63359 Al535964
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55	400045	440077 0	AA234396 N31669 H44468 AA434587 AW363088 AW993541
	100522	112277_6 19669_1	AA070906 AA070934 X51501 NM_002652 Y10179 J03460 Al791618 Al821473 AA916588 AA564296 AA916110 Al972286 Al420470 Al568790
	TOOGLE	10000_1	AI597724 AW205207 AI659305 AI791620 AA532383 AI821475 AA526498
CO	100533	32905_1	NM_012249 M31470 AL043108 AA262561 AA178863 T29433 AA313329 W48807 AW404323 AA453560 AW403227 H94816
60	100500	23902_2	W17101 AA165152 W23989 AA091310 AL121734 D54896 AA424269 BE242906 AA362118 BE018454 Al280348 AL048769 M35543 AA757734 Al128865 H20289
	100588	23902_2	H23728 Al203445 H41481 H18237 H44081 H92839 Al928621 H75675 D51148 Al796198 AW390453 D55579 D54145 D53996
			D54015 R37664 H17541 AA668681 T65061 R15867 AW468123 R16049 H69030 AA054226 H16070 F09655 R92144 T03521
<i>(</i> =			R05473 H92840 AA018186 R91707
65			U35637 AA112989 Z19308
		genbank_N62602 entrez_Z84483	
	119400	genbank_T92767	T92767
		entrez_W38197	

MISSING AT THE TIME OF PUBLICATION

TABLE 13: shows genes, including expression sequence tags, up-regulated in prostate tumor tissue compared to normal prostate tissue as analyzed using Affymetrix/Eos Hu02 GeneChip array. Shown are the ratios of "average" normal prostate to "average" prostate cancer tissues.

5

10	Pkey: ExAcon: Unigene Unigene R1:	ID:	Exemplar A Unigene nu Unigene ge		
	Pkey	ExAccn	UnigenelD	Unigene Title	R1
15	333516 337954	R73299	He 2043E4	CH22_FGENES.173_1 CH22_EM:AC005500.GENSCAN.96-3 ras homolog gene family; member B	0.028 0.029 0.03
20	337944 334111 333657	N/3289	115.204004	CH22_EM:AC005500.GENSCAN.89-7 CH22_FGENES.330_10	0.033 0.033 0.034 ~
20	327718 336355	AL137354		CH22_FGENES.241_2 CH.04_hs gi[6525284 CH22_FGENES.817_5 EST cluster (not in UniGene)	0.034 0.035 0.035
25	336377 300254 330096	AW079607	Hs.188417	CH22_FGENES.821_5 ESTs; Weakly similar to ZnT-3 [H.sapiens] CH.19_p2 gij6015278	0.036 0.037 0.037
20	335191 334040 333586			CH22_FGENES.507_6 CH22_FGENES.322_8 CH22_FGENES.204_2	0.038 0.039 0.04
30	329517	Al088120	Hs.122329	CH.10_p2 gij3983513	0.042 0.043 0.043
35	333403 335226 335976 333637			CH22_FGENES.144_21 CH22_FGENES.513_11 CH22_FGENES.652_11 CH22_FGENES.229_2	0.043 0.044 0.045 0.046
· .	334582 336437 337461			CH22_FGENES.407_5 CH22_FGENES.426_4 CH22_FGENES.782-1	0.046 0.047 0.047
40		N58545	Hs.6975	histone deacetylase 3 CH22_EM:AC005500.GENSCAN.475-3 CH22_FGENES.421_32	0.049 0.049 0.049
45	335498	AA864572 Al682088	Hs.223368	EST singleton (not in UniGene) with exon hit CH22_FGENES.571_7 ESTs	0.049 0.05 0.05
		AW025661 Al922374	Hs.240090 Hs.158549		0.051 0.052 0.052
50	335844 325371	AW247083		CH22_FGENES.54_6 EST cluster (not in UniGene) CH22_FGENES.623_4 CH.12_hs gij5866920	0.052 0.053 0.053 0.054
55	335667 333635 336736 335893			CH22_FGENES.590_18 CH22_FGENES.228_2 CH22_FGENES.110-2 CH22_FGENES.635_1	0.054 0.054 0.055 0.055
60	333170 329768 334030		Ua 497440	CH22_FGENES.94_5 CH.14_p2 gij8015501 CH22_FGENES.320_2	0.055 0.055 0.055 0.055
	300453 334262	AA234172 AW051431 Al000246	Hs.137418 Hs.113029	ESTS ribosomal protein S25 CH22_FGENES.367_12 EST singleton (not in UniGene) with exon hit	0.055 0.055 0.055 0.055
65		R22520	Hs.23398	ESTs CH22_EM:AC005500.GENSCAN.450-18 CH22_DA59H18.GENSCAN.28-5	0.055 0.056 0.056
		A1452732		EST singleton (not in UniGene) with exon hit	0.057

	339067			CH22_DA59H18.GENSCAN.33-3	0.057
	335689			CH22_FGENES.596_4	0.057
	339069			CH22_DA59H18.GENSCAN.33-5	0.057
5	338176			CH22_EM:AC005500.GENSCAN.219-4	0.057
3	328159			CH.06_hs gi[5868065	0.058
	335655			CH22_FGENES.590_6	0.058
	336371			CH22_FGENES.820_1	0.058
	336558			CH22_FGENES.842_3	0.059
10	337738			CH22_EM:AC000097.GENSCAN.100-4	0.059
10	334273 335889			CH22_FGENES.369_2	0.059
				CH22_FGENES.633_3 CH.05_hs qil5867968	0.059
	327807 333315			CH22 FGENES.138 7	0.059
	338825				0.059
15	337612			CH22_DJ246D7.GENSCAN.4-6 CH22_C20H12.GENSCAN.22-5	0.06 0.06
15	333897			CH22_FGENES.293_4	0.06
	335990			CH22_FGENES.655_4	0.06
	334264			CH22_FGENES.367_15	0.06
	338653			CH22_EM:AC005500.GENSCAN.460-39	0.061
20		W07459		EST cluster (not in UniGene)	0.061
	333498			CH22_FGENES.168_8	0.061
	336522			CH22_FGENES.839_3	0.061
		AW295677	Hs.137840	ESTs; Moderately similar to HOMEOBOX	0.00.
				PROTEIN SIX1 [H.sapiens]	0.062
25	305917	AA876469	Hs.181357	laminin receptor 1 (67kD; ribosomal protein SA)	0.062
	336143			CH22_FGENES.705_5	0.063
	333493			CH22_FGENES.168_2	0.063
	332533	M99487	Hs.1915	folate hydrolase (prostate-specific membrane antigen) 1	0.063
~~	325844			CH.16_hs gi 6552453	0.063
30	336402			CH22_FGENES.823_17	0.063
	335767			CH22_FGENES.607_1	0.064
		T80334		EST duster (not in UniGene) with exon hit	0.064
		AW177009		EST cluster (not in UniGene)	0.064
35		AA845997		EST singleton (not in UniGene) with exon hit	0.064
33	335188			CH22_FGENES.507_3	0.065
	337533			CH22_FGENES.828-2	0.065
	333311 335668			CH22_FGENES.138_3	0.065
		Al041589		CH22_FGENES.590_19 EST singleton (not in UniGene) with exon hit	0.065 0.066
40		AA962086		EST singleton (not in UniGene) with exon hit	0.066
		AA933840		EST singleton (not in UniGene) with exon hit	0.066
	335018	74 200010		CH22_FGENES.474_6	0.066
	333594			CH22_FGENES.210_3	0.066
	333900			CH22_FGENES.293_7	0.066
45	325207			CH.10_hs gi 6552430	0.067
	329888			CH.15_p2 gil6067149	0.067
	326238			CH.17_hs gij5867260	0.067
	333658			CH22_FGENES.241_4	0.067
	335809			CH22_FGENES.617_6	0.068
50	307427	Al243437		EST singleton (not in UniGene) with exon hit	0.068
	318428	Al949409	Hs.224583		0.069
	327005			CH.21_hs gi 5867664	0.069
		HG998-HT998		Sulfotransferase, Phenol-Preferring	0.069
F F	333318			CH22_FGENES.138_10	0.07
55	333313			CH22_FGENES.138_5	0.07
	325937			CH.16_hs glj5867132	0.07
	335663			CH22_FGENES.590_14	0.07
	335349	4.4004.770	11- 05 100	CH22_FGENES.539_2	0.07
60		AA224470	Hs.25426	ESTs; Weakly similar to unknown [H.sapiens]	0.07
00		N66681	Hs.33470	ESTS	0.07
	333310	A34/04/0040		CH22_FGENES.138_2	0.071
		AW340812		EST singleton (not in UniGene) with exon hit	0.071
	336340	Al453365	He 170000	CH22_FGENES.814_15 collagen; type I; alpha 1	0.071
65		AI453365 AI055966	113.172920	EST singleton (not in UniGene) with exon hit	0.071
33	335499	いいつ900		CH22_FGENES.571_8	0.071
	329669			CH22_FGENES.571_6 CH.14_p2 gij6272129	0.071 0.071
		D28390		EST duster (not in UniGene)	0.071
	338174	22000		CH22_EM:AC005500.GENSCAN.219-2	0.071
	550117				0.012

				•	
	336556			CH22_FGENES.842_1	0.072
		AA738105	Hs.140	immunoglobulin gamma 3 (Gm marker)	0.072
	336684			CH22_FGENES.46-1	0.072
5	326943			CH.21_hs gi]6004446	0.073
5	333947			CH22_FGENES.303_1	0.074
	333214		U= 474007	CH22_FGENES.104_5	0.074
		AA446572	ns.174007	ESTs; Moderately similar to !!!! ALU SUBFAMILY J WARNING	0.074
	339102 328122		•	CH22_DA59H18.GENSCAN.44-9	0.074
10		N62712	He gaeaaa	CH.06_hs gi[5868031 KIAA0618 gene product	0.075
10	328506	1102/12	115.220223	CH.07_hs gil5868471	0.075
		AA291468	Hs.98504		0.075 0.075
	335193	74201400	113.50504	CH22_FGENES.507_8	0.075
		AA971718	Hs.128141		0.076
15		AA458708		hemoglobin; alpha 2	0.076
	313644	Al565766	Hs.124960		0.076
	326145			CH.17_hs gi 5867204	0.076
	336394			CH22_FGENES.823_6	0.077
••	306516	AA989542		EST singleton (not in UniGene) with exon hit	0.077
20	300629	AA152119	Hs.155101	ATP synthase; H+ transporting; mitochondrial F1 complex; alpha s	ubunit;
				isoform 1; cardiac muscle	0.077
	333160			CH22_FGENES.91_2	0.077
	337490			CH22_FGENES.799-5	0.077
25		AA723748		EST singleton (not in UniGene) with exon hit	0.077
25		AA281765	Hs.193689		0.077
	332792	MOTOET	Un 100004	CH22_FGENES.3_2	0.078
		M81057 Al859636	Hs.8102	carboxypeptidase B1 (tissue) ribosomal protein S20	0.078
	337419	MODEOSO	113.0102	CH22_FGENES.759-4	0.078
30	333459			CH22_FGENES.157 8	0.078 0.078
•	334851			CH22_FGENES.440_3	0.078
	329046			CH.X_hs gi[5868569	0.078
	327879			CH.06_hs gi[5868142	0.079
	305830	AA857665		EST singleton (not in UniGene) with exon hit	0.079
35	302928	AL137719		EST cluster (not in UniGene) with exon hit	0.079
		AA136698	Hs.113029	ribosomal protein S25	0.079
	326390			CH.19_hs gi 5867340	0.079
	335230			CH22_FGENES.514_2	0.08
40	334622			CH22_FGENES.412_6	0.08
40	335331	AA578840	U. 77004	CH22_FGENES.535_4	80.0
		Al418863	Hs.77961	major histocompatibility complex; class I; B EST cluster (not in UniGene) with exon hit	80.0
	336561	A110000		CH22_FGENES.842_6	0.081 0.081
	335611			CH22_FGENES.583_5	0.081
45		AA635771		EST singleton (not in UniGene) with exon hit	0.081
		AA905130		EST singleton (not in UniGene) with exon hit	0.082
	308289	Al571211		EST singleton (not in UniGene) with exon hit	0.082
	334365			CH22_FGENES.378_13	0.082
~^	335496			CH22_FGENES.571_4	0.082
50	332634	S38953		Human unidentified gene complementary to P450c21	
				gene; partial cds	0.082
	337824			CH22_EM:AC005500.GENSCAN.13-18	0.082
	335822			CH22_FGENES.619_7	0.082
55	334758	ANALO 4020	H- 252400	CH22_FGENES.428_7	0.082
<i>JJ</i>	333064	AW194230	Hs.253100	CH22 FGENES.75 7	0.082
•	338695			CH22_FGENES.75_7 CH22_EM:AC005500.GENSCAN.477-25	0.083
		AA402482	Hs.97312	ESTs	0.083 0.083
	326138			CH.17_hs gi 5867203	0.083
60	328304			CH.07_hs gi 6004478	0.083
	330570	U60276	Hs.165439	arsA (bacterial) arsenite transporter; ATP-binding; homolog 1	0.083
	334305			CH22_FGENES.373_8	0.083
	335885			CH22_FGENES.632_3	0.083
~~	325839			CH.16_hs gi 6552452	0.083
65	333531			CH22_FGENES.175_18	0.084
	330385	AA449749	Hs.31386	ESTs; Highly similar to secreted apoptosis related protein	
				1 [H.sapiens]	0.084
		AA811351		Homo sapiens clone 24812 mRNA sequence	0.084
	331698	といろろとろ	Hs.65843	ESTs	0.084

	335888			CH22_FGENES.633_2		0.084
	306008	AA894390		EST singleton (not in UniGene) with exon hit		0.084
	334249			CH22_FGENES.365_15		0.084
_		AW451197	Hs.113418			0.084
5	330171			CH.02_p2 gi 6648220		0.084
	336662		11. 40-4-0	CH22_FGENES.41-1		.0.085
	320506	Al815668	HS.15/4/6	suc1-associated neurotrophic factor target 2		0.005
	040074	A1740704	U- 100000	(FGFR signalling adaptor)		0.085
10		Al740721	Hs.128292	CH22_FGENES.832_9		0.085 0.085
10	336492 335750			CH22_FGENES.602_4		0.085
	335676			CH22 FGENES.594 1		0.086
	336093			CH22 FGENES.691 2		0.086
		Al933861	Hs.222852			0.086
15	335160		***************************************	CH22_FGENES.502_4		0.086
	334306			CH22_FGENES.373_9		0.086
	334793			CH22_FGENES.433_5	•	0.086
	333936			CH22_FGENES.301_2		0.087
••	336413			CH22_FGENES.823_35		0.087
20	333775			CH22_FGENES.272_6	• ,	0.087
	335971			CH22_FGENES.652_4		0.087
		Al815981		EST cluster (not in UniGene) with exon hit		0.087
	339101			CH22_DA59H18.GENSCAN.44-6		0.087
25	327612			CH.04_hs gi 6525283		0.087 0.088
23	326241 338386			CH.17_hs gi 5867260 CH22_EM:AC005500.GENSCAN.331-4		0.088
	327762			CH.05_hs gi 5867961		0.088
		AA679772		EST singleton (not in UniGene) with exon hit		0.088
•	334359	74070772		CH22_FGENES.378_4		0.088
30	335500			CH22_FGENES.571_10		0.088
_	329687			CH.14_p2 gi 6117856		0.088
	333654			CH22_FGENES.240_2		0.088
	324430	AA464018		EST cluster (not in UniGene)		0.088
25	325999			CH.16_hs gi 5867073		0.089
35	334832			CH22_FGENES.439_1		0.089
	339115	41040000	11. 040000	CH22_DA59H18.GENSCAN.49-3		0.089
		Al916902	Hs.213882	· · · · · · · · · · · · · · · · · · ·		0.089 0.089
	328784 335044			CH.07_hs gi 5868309 CH22_FGENES.480_1		0.089
40	329791			CH.14_p2 gi[6469354		0.089
-10	333656			CH22_FGENES.240_4		0.089
	326180			CH.17_hs gi 5867211		0.089
	333391			CH22_FGENES.144_6		0.089
	338324			CH22_EM:AC005500.GENSCAN.306-3		0.089
45	305396	AA721052		EST singleton (not in UniGene) with exon hit		0.089
	337483			CH22_FGENES.795-7		0.09
	326424			CH.19_hs gi 5867369		0.09
		AA977992		EST singleton (not in UniGene) with exon hit		0.09
50	338893			CH22_DJ32I10.GENSCAN.7-6		0.09 0.09
20	327470 333165			CH.02_hs gi 5867772 CH22_FGENES.91_7		0.09
		Al186738	He 192/28	ribosomal protein S2		0.09
		AA233926	Hs.23635		•	0.09
	335334	74 200020	11022000	CH22 FGENES.535_10		0.09
55	335907			CH22_FGENES.636_2		0.09
	333885			CH22_FGENES.292_7		0.09
	331034	N51868	Hs.31965	ESTs; Moderately similar to 40S RIBOSOMAL		
				PROTEIN S20 [H.sapiens]		0.09
		AA534416	Hs.162185			0.09
60	328217			CH.06_hs glj5868096		0.091
	336068			CH22_FGENES.684_13		0.091
		AA295381	Hs.44423	ESTs		0.091
	328668 335309			CH.07_hs gij5868254 CH22_FGENES.532_2		0.091 0.091
65	335309			CH22_FGENES.532_2 CH22_EM:AC005500.GENSCAN.377-5	•	0.091
0 5		AA936892		EST singleton (not in UniGene) with exon hit		0.091
		AA639783		EST singleton (not in UniGene) with exon hit		0.091
		AA594811	Hs.119122	ribosomal protein L13a		0.091
•		AA968589	Hs.944	glucose phosphate isomerase		0.091
				•		

	323789	Al459812	Hs.170460	ESTs; Weakly similar to KIAA0990 protein [H.saplens]	0.092
	334910			CH22_FGENES.455_3	0.092
	326382			CH.19_hs gi 5867327	0.092
		AA489630	Hs.119004	KIAA0665 gene product	0.092
5	338534			CH22_EM:AC005500.GENSCAN.402-7	0.092
•	336449			CH22_FGENES.829_6	0.092
	333709			CH22_FGENES.250_24	0.092
	336559			CH22_FGENES.842_4	0.092
	333230			CH22_FGENES.107_10	0.093
10	333133			CH22_FGENES.83_9	0.093
10	334885			CH22_FGENES.451_11	0.093
		X02419	Hs.77274	plasminogen activator; urokinase	0.093
	336392	702410	110077214	CH22_FGENES.823_4	0.093
	334083			CH22_FGENES.327_38	0.093
15	325469			CH.12_hs gi 6017034	0.093
13	-	R09531	Hs.19039	ESTs	0.093
		AW500732	113.13000	EST cluster (not in UniGene) with exon hit	0.093
	334218	A11300732		CH22_FGENES.358_3	0.093
				CH22_FGENES.840_6	0.093
20	336542			CH22_FGENES.546-1	0.093
20	337151			CH22_FGENES.231_2	0.093
	333642 336863			CH22_FGENES.297-4	0.093
				CH22_FGENES.419_2	0.093
	334680			CH.18_hs gij5867297	0.093
25	326365			CH22 DJ32I10.GENSCAN.23-22	0.093
25	338952				0.094
	337539			CH22_FGENES.832-4	0.094
	333546			CH22_FGENES.180_2 CH22_FGENES.518_3	0.094
	335258			CH22_FGENES.168-19	0.094
30	336786	A1004477	Hs.237396		0.094
30		Al204177	П 3.23/33 0	CH22_FGENES.646_17	0.094
	335943			CH.06_hs gil5868165	0.094
	327918	A A070540		EST singleton (not in UniGene) with exon hit	0.094
	-	AA970548		CH22_FGENES.592_3	0.094
35	335671				0.094
33	335033			CH22_FGENES.475_11	0.094
	338277	A A E O 4 O 4 O	Ha 102024	CH22_EM:AC005500.GENSCAN.290-2 early B-cell factor	0.094
		AA504812			0.094
		AA654582	Hs.77039	ribosomal protein S3A CH22_FGENES.292_2	0.094
40	333880	Al864428	Hs.170880		0.094
40			Hs.129771		0.095
		AA648796 AA169345	115.129771	EST cluster (not in UniGene)	0.095
		AM (03343		CH22_FGENES.38_4	0.095
	332930 335368			CH22 FGENES.543_6	0.095
45		D79679	He 103/8/	ESTs; Weakly similar to Similarity with yeast gene	0.000
43	303007	R72672	115.150404	L3502.1 [C.elegans]	0.095
	000000	•		CH22_FGENES.727_3	0.095
	336223	A17070E7	Un 107727	ESTs; Weakly similar to Y38A8.1 gene product [C.elegans]	0.095
		A1767957	ns.19//3/	CH22 FGENES.648-3	0.095
50	337256	A1040002		EST singleton (not in UniGene) with exon hit	0.095
50		Al819263		CH22_FGENES.418_7	0.095
	334659			CH22_FGENES.635_3	0.095
	335895	AMI200061	Hs.4953	golgi autoantigen; golgin subfamily a; 3	0.095
		AW388061	HS.4900	CH22_FGENES.668_8	0.096
55	336010	U21260		EST cluster (not in UniGene) with exon hit	0.096
כנ	333612	U2 1200		CH22_FGENES.217_7	0.096
		AA584837		EST singleton (not in UniGene) with exon hit	0.096
		AA304037		CH22_FGENES.590_16	0.096
	335665	· A A000E00		EST singleton (not in UniGene) with exon hit	0.096
<i>6</i> 0		AA989598			0.096
60	335243			CH22_FGENES.516_4 CH22_FGENES.559_5	0.096
	335436	#14000FC	U= 101074		0.096
		Al420256	Hs.161271		0.097
	332810	AITOCOOA		CH22_FGENES.7_12 EST cincleton (not in UniGene) with even hit	0.097
25		A1735634		EST singleton (not in UniGene) with exon hit	0.097
65	335818			CH22_FGENES.618_6	0.097
	325838			CH.16_hs gi 6552452 CH22_FGENES.795-6	0.097
	337482				0.097
	336645			CH22_FGENES.26-1 CH22_FGENES.675-1	0.098
	337293			MIRET MENEOTONA.	0.030

						•
	329893			CH.15_p2 gi 6525313		0.098
	326533			CH.19_hs gi 5867441		0.098
	334905			CH22_FGENES.452_20		0.098
	306347	AA961144		EST singleton (not in UniGene) with exon hit		0.098
5	336676			CH22_FGENES.43-4		0.098
-	339166	٠,		CH22_DA59H18.GENSCAN.69-7		0.098
	335774			CH22_FGENES.607_10		0.098
	339216			CH22_FF113D11.GENSCAN.6-11		0.098
	335311			CH22_FGENES.532_4		0.098
10	329632			CH.11_p2 gi 6729060		0.098
10	328595			CH.07_hs gi[5868224		0.098
	326928			CH.21_hs gi 6456782		0.098
	-	Al079680	Hs.120770			
			H3.120//U			0.098
15		AA908508		EST singleton (not in UniGene) with exon hit		0.098
15		AA826544		EST singleton (not in UniGene) with exon hit		0.098
		T30280		EST cluster (not in UniGene)		0.099
	337553	44044000	11- 000000	CH22_C4G1.GENSCAN.2-1		0.099
		AA344069	HS.202699	neurexophilin 4		0.099
20		T08033		EST cluster (not in UniGene) with exon hit	•	
20	338981	5		CH22_DA59H18.GENSCAN.2-5		0.099
		R87365	HS.26058	ESTs; Weakly similar to p532 [H.sapiens]		0.099
	328348			CH.07_hs gi[5868383		0.099
		H49388	Hs.102082			0.099
~~		R07064		EST cluster (not in UniGene) with exon hit		0.099
25		AA608838	Hs.162681			0.099
	333227			CH22_FGENES.107_5		0.099
	316442	AA760894	Hs.153023			0.099
	326001			CH.16_hs gi 5867073		0.099
	334363			CH22_FGENES.378_11		0.099
30	338895			CH22_DJ32l10.GENSCAN.9-2		0.099
	327460			CH.02_hs gi 6004455		0.099
	332705	T59161	Hs.76293	thymosin; beta 10		0.1
	307806	Al351739		EST singleton (not in UniGene) with exon hit		0.1
	322800	F25037	Hs.225175	ESTs		0.1
35	304918	AA602697		EST singleton (not in UniGene) with exon hit		0.1
	334327			CH22_FGENES.375_4		0.1
	318359	AI097439	Hs.135548	ESTs		0.1
	326644			CH.20_hs gi 5867559		0.1
	334454			CH22_FGENES.388_3		0.1
40	327959			CH.06_hs gi[5868210		0.1
	323783	AA330586	Hs.131819			0.1
		Al955915	Hs.248038	major histocompatibility complex; class I; C		0.1
	339265			CH22_BA354I12.GENSCAN.10-3		0.1
	320576	AL049977	Hs.162209	Homo sapiens mRNA; cDNA DKFZp564C122		
45				(from clone DKFZp564C122)		0.1
	338132			CH22_EM:AC005500.GENSCAN.200-2		0.1
	333163			CH22_FGENES.91_5		0.101
	337584			CH22_C20H12.GENSCAN.5-1		0.101
		Al285535		EST singleton (not in UniGene) with exon hit		0.101
50	336969			CH22_FGENES.378-2		0.101
	327535			CH.02_hs gi 6525279		0.101
	328732			CH.07_hs gi 5868289		0.101
	336686			CH22_FGENES.46-3		0.101
	335777			CH22 FGENES.607 13		0.101
55	332944			CH22_FGENES.47_3		0.101
-	333174			CH22 FGENES.95 1		0.101
	336380			CH22_FGENES.821_8		0.101
		U60800	Hs.79089	sema domain; immunoglobulin domain (lg);		0.101
	330371	000000	113.73003	cytoplasmic domain; (semaphorin) 4D		0.101
60	221790	AA398721	Hs.186749			0.101
50	338915	TAMOUIT!	110.100743	CH22 DJ32I10.GENSCAN.12-1		0.101
						0.101
	334844			CH22_FGENES.439_24		
	336642			CH22_FGENES.23-4		0.101
65	334906			CH22_FGENES.452_21		0.101
UJ	333188	4141000000		CH22_FGENES.98_8		0.101
		AW299993		EST duster (not in UniGene) with exon hit		0.101
	329373	D.40570	11.00000	CH.X_hs gi 6682537		0.102
		R46576	Hs.23239	ESTs		0.102
	335856			CH22_FGENES.628_1		0.102

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	331888	AA431337	Hs.98017		0.102
	333154			CH22_FGENES.89_4	0.102
	335989			CH22_FGENES.655_2	0.102
		AA235602		EST singleton (not in UniGene) with exon hit	0.102
5	338016			CH22_EM:AC005500.GENSCAN.133-1	0.102
9	335190			CH22_FGENES.507_5	0.102
		T39486	Hs.6137	ESTs	0.102
	333697	100400	113.0137	CH22_FGENES.250_11	0.102
		A A000740		EST singleton (not in UniGene) with exon hit	0.102
10		AA989713			0.103
10	328734	11005040	11- 70740	CH.07_hs gi 5868289	
		Al205612	Hs.73742		0.103
	327424			CH.02_hs gi 5867751	0.103
	335872			CH22_FGENES.630_3	0.103
15	333572			CH22_FGENES.189_1	0.103
15	334774			CH22_FGENES.430_6	0.103
	338660			CH22_EM:AC005500.GENSCAN.462-1	0.103
	326713			CH.20_hs gi 5867595	0.103
	333994			CH22_FGENES.310_18	0.103
	335800		•	CH22_FGENES.613_4	0.103
20	318113	Al187943	Hs.132322	ESTs	0.103
	337278			CH22_FGENES.665-1	0.103
	336386			CH22_FGENES.822_6	0.103
	334790			CH22_FGENES.432_15	0.103
		AW505368		EST cluster (not in UniGene) with exon hit	0.104
25	336524			CH22_FGENES.839_5	0.104
	328936			CH.08 hs gil5868500	0.104
	335102			CH22 FGENES.494_7	0.104
		AA513644	Hs.222815	ESTs; Weakly similar to Wiskott-Aldrich Syndrome	
	00000			protein [H.sapiens]	0.104
30	307581	Al284415		EST singleton (not in UniGene) with exon hit	0.104
50		AW291683	Hs.226056		0.104
	335330	7111201000	110220000	CH22_FGENES.535_3	0.104
	337968			CH22_EM:AC005500.GENSCAN.103-2	0.104
	335627			CH22_FGENES.584_7	0.104
35	336274			CH22_FGENES.762_2	0.104
<i></i>	334730			CH22_FGENES.424_5	0.105
	334409	•		CH22_FGENES.383_6	0.105
	327237			CH.01_hs gi 5867544	0.105
	333321			CH22_FGENES.138_13	0.105
40		AA452366		EST cluster (not in UniGene) with exon hit	0.105
40	333738	AAAACOOO		CH22_FGENES.261_2	0.105
	338255			CH22_EM:AC005500.GENSCAN.276-3	0.105
	334282			CH22_FGENES.369_12	0.105
	330190				0.105
45		AW014249	Hs.158698	CH.05_p2 gi 6165182	0.105
73	338150	A11014243	115,150050	CH22 EM:AC005500.GENSCAN.207-2	0.105
					0.105
•	336719			CH22_FGENES.82-6	0.105
	330228			CH.05_p2 gij6013527	0.105
50	327801	075400	No. 074	CH.05_hs gij5867924	
J U		S75168	Hs.274	megakaryocyte-associated tyrosine kinase	0.105
	334972			CH22_FGENES.468_2	0.105
	335111			CH22_FGENES.494_19	0.106
	334483			CH22_FGENES.395_5	0.106
E E	328829			CH.07_hs gil5868337	0.106
55		M74299		EST cluster (not in UniGene) with exon hit	0.106
	334512			CH22_FGENES.398_10	0.106
	330024			CH.16_p2 gi 6671908	0.106
	321030	Al769930	Hs.233617	Homo sapiens (clone B3B3E13) Huntington's	
				disease candidate region	0.107
60	338410			CH22_EM:AC005500.GENSCAN.341-6	0.107
	334353			CH22_FGENES.376_5	0.107
	338276			CH22_EM:AC005500.GENSCAN.288-9	0.107
	329053			CH.X_hs gi 5868574	0.107
_]	336560			CH22_FGENES.842_5	0.107
65	332158	AA621363	Hs.112980		0.107
	336447			CH22_FGENES.829_4	0.107
	333703			CH22_FGENES.250_17	0.107
	326207			CH.17_hs gi 5867222	0.107
	333232			CH22_FGENES.108_1	0.107

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	334802			CH22_FGENES.435_1	0.107
		AA704983		EST cluster (not in UniGene) with exon hit	0.107
	338847			CH22_DJ246D7.GENSCAN.10-2	0.107
~	339407			CH22_DJ579N16.GENSCAN.1-9	0.108
5	337635			CH22_C20H12.GENSCAN.32-8	0.108
	334650			CH22_FGENES.417_17	0.108
	308511	A1687580		EST singleton (not in UniGene) with exon hit	0.108
	333392			CH22_FGENES.144_8	0.108
	325840			CH.16_hs gi]6552452	0.108
10	315044	AW205664	Hs.129568	ESTs	0.108
	333298			CH22_FGENES.133_4	0.108
	335157			CH22_FGENES.501_7	0.108
	333305			CH22_FGENES.137_2	0.108
	326379			CH.19_hs gi 5867327	0.108
15	335050	•		CH22_FGENES.482_1	0.108
	305185	AA663985	Hs.248038	major histocompatibility complex; class I; C	0.108
	335658			CH22_FGENES.590_9	0.108
		AA336609	Hs.10862		0.108
	337326			CH22_FGENES.699-6	0.108
20	339262			CH22_BA354I12.GENSCAN.9-6	0.108
		H54052	Hs.163639	ESTs; Weakly similar to INTERCELLULAR ADHESION	
				MOLECULE-1 PRECURSOR [H.sapiens]	0.109
	331792	AA398968	Hs.97548	EST	0.109
	333806			CH22_FGENES.278_2	0.109
25		AB033100		EST duster (not in UniGene)	0.109
25		AA435513	He 178170	ESTs; Weakly similar to DUAL SPECIFICITY	000
	001070	74400010	113.170170	PROTEIN PHOSPHATASE 3	0.87
	328775			CH.07_hs gij5868309	0.109
	335105			CH22_FGENES.494_10	0.109
30		Al283548	Hs.149668		0.109
50		T31940	115.148000	EST cluster (not in UniGene)	0.109
	333397	101040		CH22_FGENES.144_15	0.109
	336484			CH22_FGENES.831_3	0.109
	335507			CH22_FGENES.571_22	0.109
35	336373			CH22_FGENES.820_3	0.109
55	336188			CH22_FGENES.717_12	0.109
		AW081702	Hs.137329		0.109
		AVV061702	HS. 13/329	CH22_FGENES.506_4	0.109
	335185	AI066577		EST singleton (not in UniGene) with exon hit	0.109
40			Hs.195306		0.109
40		AI632322 AW080339	Hs.211911		0.109
			Hs.135209		0.105
		Al346359			0.11
	300212	AW135925	ns. 104002	biphenylhydrolase-like (serine hydrolase; breast epithelial	0.11
45	205675			Mucin-assoc.	0.11 0.11
43	325675			CH.14_hs gi 5867014	
	330095	A A 450064	N= 00200	CH.19_p2 gi 6015278	0.11
		AA453261	Hs.99309	ESTs	0.11
	334723			CH22_FGENES.421_34	0.11
50	333614			CH22_FGENES.217_9	0.11
50	337316	4.4005000	11-00054	CH22_FGENES.692-1	0.11
		AA635626	HS.02954	ferritin; heavy polypeptide 1	0.11
	338704			CH22_EM:AC005500.GENSCAN.480-3	0.11
	335385			CH22_FGENES.543_27 -	0.11
E E	338012			CH22_EM:AC005500.GENSCAN.128-10	0.11
55	329449			CH.Y_hs gij5868886	0.11
	338980			CH22_DA59H18.GENSCAN.2-4	0.11
	336553			CH22_FGENES.841_10	0.111
	330021			CH.16_p2 gij6671889	0.111
CO	327579			CH.03_hs gi 5867824	0.111
60	333099			CH22_FGENES.79_4	0.111
	337076	4446050	11- 405 40	CH22_FGENES.453-4	0.111
٠		AA456852	Hs.43543	suppressor of white apricot homolog 2	0.111
		Al005542	HS.180414	heat shock 70kD protein 10 (HSC71)	0.111
15		AA884409		EST singleton (not in UniGene) with exon hit	0.111
65		AA419217	Hs.15911	DKFZP586E1422 protein	0.111
	333780			CH22_FGENES.273_2	0.111
		AI702835		EST cluster (not in UniGene)	0.111
	308952	Al868157	Hs.224226		0.111
	309338	AW026946	Hs.181165	eukaryotic translation elongation factor 1 alpha 1	0.111

	329317			CH.X_hs gi 6381976	0.112
	333518			CH22_FGENES.173_3	0.112
	306982	Al127883		EST singleton (not in UniGene) with exon hit	0.112
_	336225			CH22_FGENES.728_2	0.112
5	333698			CH22_FGENES.250_12	0.112
	302173	Al417947	Hs.14068	ESTs	0.112
	335510			CH22_FGENES.571_25	0.112
	328042			CH.06_hs gi 5902482	0.112
	336512			CH22_FGENES.834_7	0.112
10	328541			CH.07_hs gi 5868486	0.112
		AW205118			0.112
		AF131846		Homo sapiens done 25028 mRNA sequence	0.112
		AF013956	Hs.123085	chromobox homolog 4 (Drosophila Pc class)	0.112
15		AA557351		ESTs; Moderately similar to MULTIFUNCTIONAL PROTEIN ADE2	
15·		Al937242	Hs.176590		0.112 0.112
		AW384710	Hs.125258		0.112
	333659			CH22_FGENES.241_5	0.113
	327510			CH.02_hs gi[6117815 CH22_FGENES.839_1	0.113
20	336520 338682			CH22_EM:AC005500.GENSCAN.472-1	0.113
20	334508			CH22_FGENES.398_6	0.113
•		T59538		EST cluster (not in UniGene)	0.113
		Al086929		EST singleton (not in UniGene) with exon hit	0.113
	336040	***************************************		CH22 FGENES.679 2	0.113
25		T23215		EST cluster (not in UniGene) with exon hit	0.113
	312011	AW294868	Hs.187226	ESTs	0.113
	335186			CH22_FGENES.506_5	0.113
	333607			CH22_FGENES.216_2	0.113
20		AA773530		EST singleton (not in UniGene) with exon hit	0.113
30	333686			CH22_FGENES.249_4	0.113
	334352	•		CH22_FGENES.376_3	0.113 0.114
	338195	•		CH22_EM:AC005500.GENSCAN.233-18 CH22_FGENES.206_2	0.114
	333588 339233			CH22_BA354I12.GENSCAN.2-3	0.114
35	337455			CH22_FGENES.777-1	0.114
		Al925108		EST singleton (not in UniGene) with exon hit	0.114
	328522			CH.07_hs gi 5868477	0.114
	323999	AI537333	Hs.252782	ESTs	0.114
	333517			CH22_FGENES.173_2	0.114
40	329935			CH.16_p2 gij6165200	0.114
	326226			CH.17_hs gi 5867230	0.114
	335890			CH22_FGENES.633_4	0.114
	336715			CH22_FGENES.77-1	0.114 0.114
45	327640			CH.04_hs gij5867890 CH22_DJ246D7.GENSCAN.7-1	0.114
73	338842	AA991487		EST singleton (not in UniGene) with exon hit	0.114
	336597	70031407		CH22_FGENES.266_1	0.114
		Y17456	Hs.227150	Homo sapiens LSFR2 gene; last exon	0.114
		AA159213	Hs.5337	isocitrate dehydrogenase 2 (NADP+); mitochondrial	0.114
50		N44238	Hs.77515	inositol 1;4;5-triphosphate receptor; type 3	0.114
	327358			CH.01_hs gi 6552411	0.114
	308792	Al815153	Hs.195188	glyceraldehyde-3-phosphate dehydrogenase	0.115
	325886			CH.16_hs gi 5867087 -	0.115
E E	336850			CH22_FGENES.272-11	0.115
55		AA863103		EST singleton (not in UniGene) with exon hit	0.115 0.115
		AC004472		multiple UniGene matches CH22_FGENES.707_2	0.115
	336158			CH.06_hs gij5868131	0.115
	327866 339157			CH22_DA59H18.GENSCAN.67-3	0.115
60	339258			CH22_BA354I12.GENSCAN.8-3	0.115
00	336129			CH22_FGENES.701_17	0.115
	333684	•		CH22_FGENES.249_2	0.115
		AW190162		ribosomal protein L23a	0.115
		AA954097	Hs.127523		0.115
65	302640	AB035698		EST cluster (not in UniGene) with exon hit	0.115
	328968			CH.08_hs gil6456775	0.115
	327902			CH.06_hs gi 5868158	0.115
		AJ223366		EST duster (not in UniGene)	0.115 0.115
	335962			CH22_FGENES.651_4	0.115

	334927			CH22_FGENES.460_1	0.115
	330535	U11872		Human interleukin-8 receptor type B (IL8RB) mRNA,	
				splice variant IL8RB1	0.856
	328591			CH.07_hs gi 5868227	0.115
5	334902			CH22_FGENES.452_16	0.115
-	328525			CH.07_hs gi 5868482	0.115
					0.116
	325870			CH.16_hs gi 6682492	
	337522			CH22_FGENES.819-1	0.116
10	305079	AA641329		EST singleton (not in UniGene) with exon hit	0.116
10	327343			CH.01_hs gij6017017	0.116
	333918			CH22_FGENES.296_7	0.116
	333600			CH22 FGENES.213_2	0.116
	335846			CH22_FGENES.623_6	0.116
	333510			CH22 FGENES.171_4	0.116
15	327629			CH.04_hs gi 5867872	0.116
	333470			CH22_FGENES.161_6	0.116
	326855			CH.20_hs gi 6552460	0.116
					0.117
	327008			CH.21_hs gi 5867664	0.117
20	337480			CH22_FGENES.795-3	
20	336425			CH22_FGENES.824_10	0.117
	321964	AL079687	Hs.171065		0.117
	335651			CH22_FGENES.590_2	0.117
	308164	AJ521574	Hs.181165	eukaryotic translation elongation factor 1 alpha 1	0.117
	337927			CH22 EM:AC005500.GENSCAN.80-3	0.117
25		H45095	Hs.153524	· · · ·	0.117
		Al245127	Hs.179331		0.117
	-	AA937331	110.170001	EST singleton (not in UniGene) with exon hit	0.117
		Mastrati		CH.14_p2 gi 6272129	0.117
	329670				0.117
20	335612	41000450		CH22_FGENES.583_6	
30		Al363450		EST singleton (not in UniGene) with exon hit	0.117
	330401	D28383		Human mRNA for ATP synthase B chain, 5'UTR (sequence from the	
				5'cap to the start codon)	0.117
	327127			CH.21_hs gi 6682520	0.117
	333843			CH22_FGENES.290_1	0.117
35	331083	R17762	Hs.22292	ESTs	0.117
	329140			CH.X_hs gi]6017060	0.117
	339338			CH22_BA354112.GENSCAN 27-3	0.117
		AA464518	Hs.99616	ESTs	0.117
	338631	7411010	110100010	CH22_EM:AC005500.GENSCAN.454-2	0.117
40	330299			CH.06_p2 gi 2905881	0.117
TU					0.117
	330351	4 474574 4	U- 4040F7	CH.09_p2 gi 3056622	0.117
		AA715714	HS. 16 1357	laminin receptor 1 (67kD; ribosomal protein SA)	
	333106			CH22_FGENES.79_12	0.117
45	338514			CH22_EM:AC005500.GENSCAN.392-4	0.117
45	327335			CH.01_hs gi 5902477	0.117
	301970	AB028962	Hs.120245	KIAA1039 protein	0.118
	326339			CH.17_hs gi[6056311	0.118
	330612	X15673	Hs.93174	Human endogenous retrovirus pHE.1 (ERV9)	0.118
	334178			CH22_FGENES.350_6	0.118
50	328008			CH.06_hs gi 5902482	0.118
	329976			CH.16_p2 gi 4878063	0.118
		AA897432	Hs.130411		0.118
			113.100711	EST singleton (not in UniGene) with exon hit	0.118
		AA789095		CH22_EM:AC005500.GENSCAN.34-3	0.118
55	337850				0.118
<i>33</i>	333626			CH22_FGENES.224_2	
	337672			CH22_EM:AC000097.GENSCAN.67-1	0.118
	328803			CH.07_hs gi 6004475	0.118
	325922			CH.16_hs gi 5867122	0.118
	334489			CH22_FGENES.397_1	0.118
60	320638	R54766	Hs.101120	ESTs	0.118
	321932	AA569229		EST cluster (not in UniGene)	0.118
	336958			CH22_FGENES.367-1	0.118
		AA600176	Hs.112345		0.118
		AA889992		EST singleton (not in UniGene) with exon hit	0.118
65		PARTITION		CH22 FGENES.194-1	0.118
0 3	336803	AIOOEOOO		EST singleton (not in UniGene) with exon hit	0.118
		Al925823			0.118
	336859			CH22_FGENES.293-9	
	337935	•		CH22_EM:AC005500.GENSCAN.85-6	0.118
	326492		•	CH.19_hs gi 5867422	0.118

	327289			CH.01_hs gi 5867481	0.119
	325818			CH.14_hs gi 6682490	0.119
	310787	AW262580	Hs.159040		0.119
_	330028			CH.16_p2 gi]6671908	0.119
5	325317	•		CH.11_hs gi 5866878	0.119
	335279		•	CH22_FGENES.523_7	0.119
	331720	AA192173	Hs.221530	and the second s	0.119
	329186			CH.X_hs gi 5868711	0.119
10	316012	AA764950	Hs.119898		0.119
10	338316			CH22_EM:AC005500.GENSCAN.304-2	0.119
	326033			CH.17_hs gi 5867178	0.119
	334745			CH22_FGENES.426_3	0.119
	333051			CH22_FGENES.73_5	0.119
1.5		R01279		EST cluster (not in UniGene) with exon hit	0.12
15		AA454809	Hs.172928	collagen; type I; alpha 1	0.12
	335680			CH22_FGENES.594_5	0.12
		AA548556		EST singleton (not in UniGene) with exon hit	0.12
	335441			CH22_FGENES.560_4	0.12
20	336187	****		CH22_FGENES.717_11	0.12 0.12
20		AW087175		EST singleton (not in UniGene) with exon hit	0.12
	336047	414405050		CH22_FGENES.679_9	0.12
		AW195850	II- 001000	EST singleton (not in UniGene) with exon hit	0.12
		A1695385	Hs.201903		0.12
25		AA399444		EST singleton (not in UniGene) with exon hit CH22_FGENES.746_3	0.12
23	336245	H72333		EST duster (not in UniGene) with exon hit	0.12
	335690	H/2333		CH22_FGENES.596_5	0.12
	328941			CH.08_hs gi[6456765	0.12
	333873			CH22_FGENES.291_9	0.12
30		AW105092	Hs.155690		0.12
50	339288	7111100002	. 10. 100000	CH22_BA354I12.GENSCAN.16-6	0.12
	337996			CH22 EM:AC005500.GENSCAN.116-3	0.12
	333304			CH22_FGENES.137_1	0.121
		Al591235		EST singleton (not in UniGene) with exon hit	0.121
35	329319			CH.X_hs gi]6381976	0.121
		X57138		multiple UniGene matches	0.121
	333290			CH22_FGENES.129_2	0.121
	323825	Al793080	Hs.123525	ESTs; Weakly similar to NEUTROPHIL GELATINASE-ASSOCIATE	D
4.0				LIPOCALIN PRECURSOR [R.norvegicus]	0.121
40		U64105		Rho guanine nucleotide exchange factor (GEF) 1	0.121
		AA679990	HS.181165	eukaryotic translation elongation factor 1 alpha 1	0.121
	333647	4.40000.40		CH22_FGENES.235_2	0.121 0.121
		AA333340		EST cluster (not in UniGene) with exon hit	0.121
15	329777			CH.14_p2 gi[6002090	0.121
45	333155			CH22_FGENES.89_5	0.121
	326122			CH:17_hs gi 5867194 CH:22_FGENES.532_3	0.121
	335310			CH22_FGENES.562_13	0.122
	335453	AA643329	He 11133/	ferritin; light polypeptide	0.122
50	337284	Anonoco	113.111007	CH22_FGENES.667-2	0.122
50	337418			CH22_FGENES.758-4	0.122
	313073	Al963740	Hs.46826	ESTs	0.122
		AW504164	110.40020	EST cluster (not in UniGene) with exon hit	0.122
	300017	7111001101			
55		M33197		AFFX control: GAPDH	0.122
	316725	AW135084	Hs.127264	ESTs	0.122
		AA293153	Hs.120980	nuclear receptor co-repressor 2	0.122
	336466			CH22_FGENES.829_25	0.122
	335956			CH22_FGENES.647_3	0.122
60	315308	AA780564	Hs.189053	ESTs	0.122
	338925			CH22_DJ32I10.GENSCAN.14-3	0.122
	334969			CH22_FGENES.466_2	0.122
	322050	AL137589		EST cluster (not in UniGene)	0.122
	339084			CH22_DA59H18.GENSCAN.38-2	0.122
65	338323			CH22_EM:AC005500.GENSCAN.306-2	0.122
	337003			CH22_FGENES.419-7	0.122
	325470			CH.12_hs gi 6017034	0.123
	336503	D00071	U. APAC.	CH22_FGENES.833_10	0.123
	330786	D60374	Hs.258712	ESI	0.123

	329446			CH.Y_hs gi 5868886	0.123
		AA229433	Hs.222634	ESTs; Moderately similar to ubiquitin-like protein /	
				ribosomal protein S30	0.123
	309067	Al916313	Hs.212788		0.123
5	317464	AA968472	Hs.130463	ESTs	0.123
	328755			CH.07_hs gi 5868301	0.123
	326036			CH.17_hs gi 5867178	0.123
	327208			CH.01_hs gi 5867447	0.123
	326124			CH.17_hs gi 5916395	0.123
10	327509			CH.02_hs gi 6117815	0.123
	338398			CH22_EM:AC005500.GENSCAN.336-5	0.123
	304652	AA527782	Hs.84298		
				histocompatibility complex; class II antigen-associated)	0.123
1 =	335797			CH22_FGENES.612_6	0.124
15	336714			CH22_FGENES.76-29	0.124
	327204	44400000	11. 400	CH.01_hs gij5867447	0.124
		AA430672	Hs.123778		0.124
		Al126509		EST singleton (not in UniGene) with exon hit	0.124 0.124
20	336174			CH22_FGENES.710_1	0.124
20	336126			CH22_FGENES.701_13	0.124
	329129	AMANTERS		CH.X_hs gi 6588026 EST cluster (not in UniGene) with exon hit	0.124
	335778	AW407562		CH22_FGENES.607_14	0.124
	336601			CH22_FGENES.369_2	0.124
25	334340			CH22_FGENES.375_17	0.124
24	337436			CH22_FGENES.767-1	0.124
		AA896990		EST singleton (not in UniGene) with exon hit	0.124
	339213	70 000000		CH22_FF113D11.GENSCAN.6-8	0.124
	335355			CH22_FGENES.541_2	0.124
30	336552			CH22_FGENES.841_9	0.124
	336384			CH22_FGENES.822_4	0.124
	310485	Al286202	Hs.149800	ESTs	0.125
	335840			CH22_FGENES.622_3	0.125
	336444			CH22_FGENES.827_10	0.125
35	315703	N36070		EST cluster (not in UniGene)	0.125
	327763			CH.05_hs gi 5867961	0.125
	336383			CH22_FGENES.822_3	0.125
	333496			CH22_FGENES.168_6	0.125
40	328662			CH.07_hs gi 6004473	0.125
40	338986			CH22_DA59H18.GENSCAN.5-1	0.125 0.125
	328311		•	CH.07_hs gij5868371	0.125
	337241 336933			CH22_FGENES.644-2 CH22_FGENES.350-7	0.125
		AW294432	Hs.144252		0.125
45	326116	MICOMOL	110,177606	CH.17_hs gi 5867193	0.125
		HG363-HT363		Epidermal Growth Factor Receptor-Related Protein	0.125
		Al268539		EST singleton (not in UniGene) with exon hit	0.125
• .		AA418988	Hs.98314	Homo sapiens mRNA; cDNA DKFZp586L0120	
				(from clone DKFZp586L0120)	0.125
50	330462	HG944-HT944		Dopamine Receptor D4	0.125
	304410	AA284508		EST singleton (not in UniGene) with exon hit	0.125
	336385			CH22_FGENES.822_5	0.125
	336793			CH22_FGENES.176-3	0.125
	326243			CH.17_hs gi 5867261	0.125
55	327266			CH.01_hs gi 5867462	0.125
		AF070579	Hs.181544	Homo sapiens clone 24487 mRNA sequence	0.125
	336960			CH22_FGENES.369-5	0.125
	329667			CH.14_p2 gi 6272129	0.125
<i>(</i> 0	328168			CH.06_hs gi 5868071	0.125
60	336534			CH22_FGENES.839_16	0.125
	339289			CH22_BA354112.GENSCAN.16-9	0.126
		Al970747		EST singleton (not in UniGene) with exon hit	0.126
	339190			CH22_FF113D11.GENSCAN.1-2	0.126 0.126
65	337086	D01054	Hs.211522	CH22_FGENES.458-14	0.126
00		R21054	113.611362	ES18 CH22_BA232E17.GENSCAN.6-8	0.126
	339396	A A / / O O 7 7	Hs 170765	Homo sapiens mRNA; cDNA DKFZp586H1921	V.120
	331830	AA449077	.10.11.07.03	(from clone DKFZp586H192	0.126
	302000	A1475914		EST singleton (not in UniGene) with exon hit	0.126
	000000	, 4410014		Baton from at outdoing that over the	

	338477			CH22_EM:AC005500.GENSCAN.373-5	0.126
	334286			CH22_FGENES.369_16	0.126
		A1025039	Hs.131732		0.126
		7400000	113.101762	CH22_FGENES.516_10	0.126
_	335249				
5	333327			CH22_FGENES.138_20	0.126
	304240	AA009802		EST singleton (not in UniGene) with exon hit	0.126
	335464			CH22_FGENES.562_26	0.126
	335236			CH22_FGENES.515_8	0.126
					0.126
10	334154			CH22_FGENES.340_4	
10	309257	Al984183		EST singleton (not in UniGene) with exon hit	0.126
	310015	Al220122	Hs.201981	ESTs; Weakly similar to breast carcinoma-associated antigen	
•				[H.sapiens]	0.126
	328280	•			0.126
		4.4004040		CH.07_hs gi 5868352	
		AA831819		EST singleton (not in UniGene) with exon hit	0.126
15	327430			CH.02_hs gi 5867754	0.126
	328323			CH.07_hs gi[5868373	0.126
	333274			CH22_FGENES.123_2	0.126
					0.127
	337193			CH22_FGENES.575-3	
	334820			CH22_FGENES.437_2	0.127
20	328706			CH.07_hs gi 5868270	0.127
		W67267	Hs.174911		0.127
		Al192479		EST singleton (not in UniGene) with exon hit	0.127
		A1192413			
	337123			CH22_FGENES.519-3	0.127
	326201			CH.17_hs gi 5867216	0.127
25	335276			CH22_FGENES.523_2	0.127
	331202	T81115	Hs.191136		0.127
		U03187			0.127
			NS. 12 (344	interleukin 12 receptor, beta 1	
		N49521		EST duster (not in UniGene)	0.127
	301743	F12605	Hs.204529	ESTs; Weakly similar to reverse transcriptase [H.sapiens]	0.127
30	328175			CH.06_hs gi[5868073	0.127
	306407	AA971985		EST singleton (not in UniGene) with exon hit	0.127
		70007 1000			0.127
	327145			CH.01_hs gi 5867548	
	327649			CH.04_hs gi[5867899	0.127
	335142			CH22_FGENES.498_12	0.127
35	333909			CH22_FGENES.295_2	0.127
-		X04325	Hs.2679	gap junction protein; beta 1; 32kD (connexin 32;	
	330000	AU-1020	113.2013		0.127
				Charcot-Marie-Tooth neuropathy; X-linked)	
	330158			CH.21_p2 gi 6580367	0.127
	320153	AF064594	Hs.120360	phospholipase A2; group VI	0.127
40	314407	AA098835	Hs.224432	ESTs	0.127
	333383			CH22_FGENES.143_22	0.127
		A 170 40 40	11-044470		0.128
		A1734242	Hs.244473		
	326233			CH.17_hs gi 5867232	0.128
	326598			CH.20_hs gij5867634	0.128
45	335174			CH22_FGENES.504_4	0.128
		H29920	Hs.99486	ESTs; Weakly similar to aralar1 [H.sapiens]	0.128
		1123320	115.55400		
	335458	•		CH22_FGENES.562_18	0.128
	332997			CH22_FGENES.58_4	0.128
	334188			CH22_FGENES.352_3	0.128
50	329759			CH.14_p2 gi 6048280	0.128
50				CH.09_p2 gi 4544475	0.128
	330348			011.03_pz gij4044473	
	326958			CH.21_hs gi 6469836	0.128
	305263	AA679467		EST singleton (not in UniGene) with exon hit	0.128
	337693			CH22 EM:AC000097.GENSCAN.78-14	0.128
55	326812			CH.20_hs gi[6682504	0.128
55				CH22_FGENES.108_7	0.128
	333237				
	333699			CH22_FGENES.250_13	0.128
	311496	Al768677	Hs.209888	ESTs; Weakly similar to phosphatidylserine	
		•		synthase-2 [M.musculus]	0.128
60	336499			CH22_FGENES.833_4	0.128
50		A Engagory	Un 44000F		0.128
		AF032387		small nuclear RNA activating complex; polypeptide 4; 190kD	
	309989	Al184186	Hs.197813	ESTS	0.128
	301490	AW298468	Hs.250461	ESTs	0.128
	337011			CH22_FGENES.427-6	0.128
65		A A 976010	Hs.134427	. -	0.128
U.J		AA876910			
		W22172	Hs.59038		0.128
	336497			CH22_FGENES.833_2	0.129
				and the state of the second section and the second section of	A 400
	302068	Y16280	Hs.132049	endomelin type o receptor-like protein 2	0.129
	302068 334502	Y16280	Hs.132049	endothelin type b receptor-like protein 2 CH22_FGENES.397_18	0.129

	304332	AA158884		EST singleton (not in UniGene) with exon hit	0.129
	304522	AA465405		EST singleton (not in UniGene) with exon hit	0.129
		R46180	Hs.153485		0.129
			Hs.161354		
~		Al685841	MS. 10 1354		0.129
5	301119	AF142579		EST duster (not in UniGene) with exon hit	0.129
	309268	Al985821	Hs.62954	ferritin; heavy polypeptide 1	0.129
		H42142		DEAD/H (Asp-Glu-Ala-Asp/His) box polypeptide 19	
	000000	1176176		(Dbp5; yeast; homolog)	0.129
	000040			3	
10	336949		•	CH22_FGENES.361-4	0.129
10	330115			CH.19_p2 gi 6015202	0.129
	339212			CH22_FF113D11.GENSCAN.6-7	0.129
	326951			CH.21_hs gij6004446	0.129
		AA662939		EST singleton (not in UniGene) with exon hit	0.129
1 ~		Al559492		EST singleton (not in UniGene) with exon hit	0.129
15	337140			CH22_FGENES.537-5	0.13
	321758	U29112		EST duster (not in UniGene)	0.13
	304619	AA515554	Hs 119598	ribosomal protein L3	0.13
		AA745289	Hs.173088		0.13
		AA140203	115.115000		0.13
00	339017			CH22_DA59H18.GENSCAN.20-6	
20	330116			CH.19_p2 gi 6015202	0.13
	333312	•		CH22_FGENES.138_4	0.13
	338004	•		CH22_EM:AC005500.GENSCAN.121-1	0.13
		AA232134	Hs.190028	-	0.13
					0.13
25		A1239845	∏5.120494	ESTs; Weakly similar to EG:9587.2 [D.melanogaster]	
25	338530			CH22_EM:AC005500.GENSCAN.398-11	0.13
	335968			CH22_FGENES.652_1	0.13
	314121	Al732100	Hs.187619	ESTs	0.13
	337593			CH22_C20H12.GENSCAN.6-8	0.13
	332881			CH22_FGENES.33_1	0.13
30		A A050040			0.13
30		AA858043		EST singleton (not in UniGene) with exon hit	
	339059			CH22_DA59H18.GENSCAN.30-5	0.13
	305610	AA782319		EST singleton (not in UniGene) with exon hit	0.13
	305852	AA862455		EST singleton (not in UniGene) with exon hit	0.13
	327409			CH.02_hs gl 5867750	0.13
35		A1649090	Hs.164178		0.13
33		Al613089			
		Al799268	Hs.209929		0.13
	325961			CH.16_hs gi 5867147	0.13
	311159	AW025919	Hs.197636	ESTs	0.13
	322715	AA057230	Hs.182135	ESTs	0.13
40	336441			CH22_FGENES.827_7	0.13
10	336339			CH22_FGENES.814_12	0.13
		41005005			
		Al095365		EST singleton (not in UniGene) with exon hit	0.13
	333613			CH22_FGENES.217_8	0.13
	338489			CH22_EM:AC005500.GENSCAN.384-17	0.131
45	326904			CH.21_hs gi 5867684	0.131
	337337			CH22_FGENES.717-1	0.131
	326752			CH.20_hs gi 5867615	0.131
		111/5/0070		CH20_its yipoorots	
		AW512978		EST singleton (not in UniGene) with exon hit	0.131
	301373	AA595235		EST cluster (not in UniGene) with exon hit	0.131
50	338448			CH22_EM:AC005500.GENSCAN.359-22	0.131
	333774			CH22_FGENES.272_5	0.131
	332986			CH22_FGENES.54_8	0.131
	335362				0.131
				CH22_FGENES.541_12 -	
	335896			CH22_FGENES.635_4	0.131
55	337825			CH22_EM:AC005500.GENSCAN.13-19	0.131
	325257			CH.11_hs gi 5866895	0.131
		T50240	Hs.167837	FSTs	0.131
		Y08302		dual specificity phosphatase 9	0.131
CO		AA292721		ESTs; Weakly similar to unknown [H.saplens]	0.131
60	322995	AA513829	Hs.29797	ribosomal protein L10	0.131
	335497			CH22_FGENES.571_5	0.131
	334824			CH22_FGENES.437_6	0.131
		R06933	Hs.184221		0.131
		1100300	113.104221		
CF	334842			CH22_FGENES.439_21	0.131
65	333335			CH22_FGENES.139_4	0.131
	317252	AA905178	Hs.130124	ESTs	0.131
	329034			CH.X hs ail5868561	0.131
		AA664230		EST singleton (not in UniGene) with exon hit	0.131
		TOUTEUU			
	335755			CH22_FGENES.604_4	0.131

	302143	H15270	Hs.189847	putative neuronal cell adhesion molecule	0.131
	334939			CH22_FGENES.465 3	0.131
		C15110	Hs.17802		0.131
		Ciario	115.17002		
	334498			CH22_FGENES.397_14	0.131
5	333413			CH22_FGENES.146_2	0.132
	329676			CH.14_p2 gi[6272128	0.132
	327277			CH.01_hs gij5867473	0.132
		AA627416		EST singleton (not in UniGene) with exon hit	0.132
		741027410		OLION ECCINES 400 0	
10	336805			CH22_FGENES.196-3	0.132
10	320121	T93657		EST cluster (not in UniGene)	0.132
	334761			CH22_FGENES.428_10	0.132
	339400			CH22 BA232E17.GENSCAN.7-6	0.132
	330301			CH.06_p2 gi 2905862	0.132
		A A007004	No 100007		U. 10L
15	310022	AA827691	ris.12880/	ESTs; Weakly similar to neuronal thread protein	0.400
15				AD7c-NTP [H.sapiens]	0.132
	328020			CH.06_hs gi 5902482	0.132
	325327			CH.11_hs gij5866875	0.132
		AA209530		EST cluster (not in UniGene)	0.132
	336393			CH22_FGENES.823_5	0.132
20					
20	325905			CH.16_hs gi 5867104	0.132
	305237	AA676286	Hs.2186	eukaryotic translation elongation factor 1 gamma	0.132
	339046			CH22_DA59H18.GENSCAN.28-6	0.132
	325375			CH.12_hs gi 5866920	0.132
	333961			CH22_FGENES.304_7	0.132
25				CH22_FGENES.562_8	0.133
23	335450	D00400			
		R58438		EST duster (not in UniGene) with exon hit	0.133
	335116			CH22_FGENES.496_3	0.133
	327333			CH.01_hs gi 5902477	0.133
	308070	Al470948		EST singleton (not in UniGene) with exon hit	0.133
30		AI581855		EST singleton (not in UniGene) with exon hit	0.133
50			Hs.208839		0.133
		AW360847	115.200005		
		AW248307		EST cluster (not in UniGene)	0.133
	328318			CH.07_hs gi 5868373	0.133
	320603	R51419		EST cluster (not in UniGene)	0.133
35	332791			CH22_FGENES.3_1	0.133
-		AA524725	Hs.162108		0.133
			Hs.224868		0.133
		AL134164			
		R39753	Hs.170187		0.133
40	333944			CH22_FGENES.302_2	0.133
40	317992	Al733512	Hs.130901	ESTs	0.133
	330935	F02383	Hs.26492	beta-1;3-glucuronyltransferase 3 (glucuronosyltransferase i)	0.133
	336659			CH22_FGENES.36-5	0.133
	338887			CH22_DJ32I10.GENSCAN.6-10	0.133
		4.4.070.070	U= 40440E		
45		AA679979	IDS. 18 1 100	eukaryotic translation elongation factor 1 alpha 1	0.133
45	333566			CH22_FGENES.183_2	0.134
	316952	AW450033	Hs.163312	ESTs	0.134
	333818			CH22_FGENES.283_1	0.134
	328687			CH.07_hs gi 5868262	0.134
		H11802		EST cluster (not in UniGene) with exon hit	0.134
50		1111002			0.134
J U	336557			CH22_FGENES.842_2	
	335222			CH22_FGENES.513_5	0.134
	338094			CH22_EM:AC005500.GENSCAN.179-3	0.134
	337384			CH22_FGENES.745-1 -	0.134
	327360			CH.01 hs gil6552411	0.134
55	328132			CH.06_hs gi 5868038	0.134
33		A1751490	Uc 192927	ESTs; Weakly similar to !!!! ALU SUBFAMILY SQ	0.101
	323004	Al751438	115.102021		0.404
				WARNING ENTRY !!!!	0.134
	337591			CH22_C20H12.GENSCAN.6-6	0.134
	307018	Al140639		EST singleton (not in UniGene) with exon hit	0.134
60	326896			CH.21_hs gij5867680	0.134
	333479			CH22_FGENES.163_5	0.134
	337915			CH22_EM:AC005500.GENSCAN.61-3	0.134
•	335110			CH22_FGENES.494_18	0.134
	333481			CH22_FGENES.163_9	0.134
65	327512			CH.02_hs gl 6117815	0.134
		AW328639	Hs.83575	ESTs; Weakly similar to ZC328.3 [C.elegans]	0.134
	330163			CH.02_p2 gij6042042	0.135
				CH22_FGENES.604_1	0.135
	335752				
	334857			CH22_FGENES.443_1	0.135

	301872	H84730		EST cluster (not in UniGene) with exon hit	0.135
	337529			CH22_FGENES.823-29	0.135
	335734			CH22_FGENES.601_4	0.135
_	337551			CH22_FGENES.847-8	0.135
5		Al920965	Hs.77961	major histocompatibility complex; class I; B	0.135
	335513			CH22_FGENES.571_28	0.135
	339078			CH22_DA59H18.GENSCAN.37-6	0.135
	321907	N56660	Hs.148722	ESTs; Weakly similar to large tumor suppressor 1 [H.sapiens]	0.135
	337189			CH22_FGENES.571-32	0.135
10	329635			CH.12_p2 gi 5302817	0.135
10		Al719930		EST singleton (not in UniGene) with exon hit	0.135
			11- 0004	· · · · · · · · · · · · · · · · · · ·	
		AA627248	Hs.2064	vimentin	0.135
	333894			CH22_FGENES.293_1	0.135
	322465	AA137152	Hs.3784	ESTs; Highly similar to phosphoserine aminotransferase	
15				[H.sapiens]	0.135
	305601	AA780975		EST singleton (not in UniGene) with exon hit	0.135
	332186	H10781	Hs.141051	ESTs; Moderately similar to IIII ALU SUBFAMILY SB	
				WARNING ENTRY	0.135
	327822			CH.05_hs gi 5867968	0.135
20		Al393914	Un 160694	ESTs; Weakly similar to similar to CR16; SH3 domain	0
20	310001	MI0303 14	F15.100024		0.195
				binding protein	0.135
	328752			CH.07_hs gi 5868298	0.135
	337611			CH22_C20H12.GENSCAN.19-4	0.135
	334470			CH22_FGENES.394_1	0.136
25	335115			CH22 FGENES.496 2	0.136
	328730			CH.07_hs gt 5868289	0.136
	330350			CH.09_p2 gi[3056622	0.136
	336971			CH22 FGENES.378-6	0.136
		AIECEC10		EST singleton (not in UniGene) with exon hit	0.136
30		Al565612			
20	326745			CH.20_hs gi 5867611	0.136
	335440			CH22_FGENES.560_3	0.136
	320257	AA330746		EST cluster (not in UniGene)	0.136
	328677			CH.07_hs gi 5868256	0.136
	329731			CH.14_p2 gij6065783	0.136
35	315950	AA700553	Hs.206974		0.136
	330049			CH.17_p2 gi 4567182	0.136
	337070			CH22_FGENES.448-3	0.136
		H11324	Hs.31059		0.136
					0.136
40		AW005527	Hs.232820		
40	333458			CH22_FGENES.157_7	0.136
	329899			CH.15_p2 gi 6563505	0.136
	322202	Al275056	Hs.200133	ESTs	0.136
	333991			CH22_FGENES.310_15	0.136
	318617	AW247252	Hs.75514	nucleoside phosphorylase	0.136
45	310623	Al341586	Hs.195588	ESTs	0.136
-		M23323	Hs.3003	CD3E antigen; epsilon polypeptide (TiT3 complex)	0.136
		AW194694		EST singleton (not in UniGene) with exon hit	0.136
		R00071	Hs.191199	• • •	0.136
		noor I	110.101100	CH22_FGENES.369_15	0.136
50	334285	E40000	11. 400705	-	
50		F13689	Hs.100725		0.136
		AA827608		EST singleton (not in UniGene) with exon hit	0.136
	303158	AL138110	Hs.8594	Homo sapiens mRNA containing (CAG)4 repeat; clone CZ-CAG-7	
	334543			CH22_FGENES.403_8 -	0.136
	335384			CH22_FGENES.543_26	0.136
55	336527			CH22_FGENES.839_8	0.136
	334951			CH22_FGENES.465_20	0.136
	325882			CH.16_hs gij5867087	0.137
		AA653159		EST singleton (not in UniGene) with exon hit	0.137
60		A1148709	Un danner	EST singleton (not in UniGene) with exon hit	0.137
60		AA453418	Hs.178272		0.137
		R44780	Hs.22634	ESTs	0.137
	306094	AA908877		EST singleton (not in UniGene) with exon hit	0.137
	333561			CH22_FGENES.180_18	0.137
		H61962		EST duster (not in UniGene)	0.137
65		AA497090		EST cluster (not in UniGene)	0.137
	337926			CH22_EM:AC005500.GENSCAN.77-4	0.137
	337353			CH22_FGENES.726-1	0.137
		AA41000E	Hs.104774		0.137
		AA412295	ns.104774		
	308981	A1873242		EST singleton (not in UniGene) with exon hit	0.137

	000404			OUV ha affected to	0.107
	329424			CH.Y_hs gi[5868879	0.137
	325829	4.4.4.0000	11- 00400	CH.15_hs gi 5867052	0.137
		AA416863	Hs.98183	ESTS	0.137
5	333854	41000040		CH22_FGENES.290_13	0.137
5		A1000248		EST singleton (not in UniGene) with exon hit	0.137
	328948			CH.08_hs gi 6456765	0.137
	338935			CH22_DJ32i10.GENSCAN.18-12	0.137 0.137
	325960			CH.16_hs gi 5867147	0.137
10	328377	Algongon		CH.07_hs gij5868390 EST singleton (not in UniGene) with exon hit	0.138
10		Al829820 AA424352	Hs.210586		0.138
	337592	MMZ400Z	115.2 10000	CH22_C20H12.GENSCAN.6-7	0.138
	338684			CH22_EM:AC005500.GENSCAN.472-3	0.138
		AA400498	Hs.97543		0.138
15		AA505535	113.375-0	EST singleton (not in UniGene) with exon hit	0.138
1.5	333981	A100000		CH22_FGENES.310_4	0.138
		AA040369	Hs 11170	SYT interacting protein	0.138
		AA835278	110.11110	EST singleton (not in UniGene) with exon hit	0.138
		T65554	Hs.251591		0.138
20	333783	100007		CH22_FGENES.273_5	0.138
20	337406			CH22_FGENES.754-14	0.138
	327976		•	CH.06_hs gi 5868212	0.138
	325593			CH.13_hs gi 5866992	0.138
	339425			CH22_DJ579N16.GENSCAN.14-4	0.138
25		AA428879		EST singleton (not in UniGene) with exon hit	0.138
		AW131104		EST singleton (not in UniGene) with exon hit	0.138
	337532			CH22_FGENES.827-6	0.138
		AA904448	Hs,126368		0.138
		AA854425	Hs.144455	ESTs	0.138
30	328927			CH.08_hs gi 5868500	0.138
	336424			CH22_FGENES.824_9	0.138
	326667			CH.20_hs gi 6552455	0.138
	325988			CH.16_hs gi 5867064	0.138
~~	318446	AW300287		EST duster (not in UniGene)	0.139
35	336511			CH22_FGENES.834_6	0.139
	335204			CH22_FGENES.508_13	0.139
		AA147472		EST cluster (not in UniGene) with exon hit	0.139
		AA115804	Hs.187593		0.139
40	329376	4.4500000		CH.X_hs gi 5868859	0.139 0.139
40		AA563898		EST singleton (not in UniGene) with exon hit CH22_FGENES.239_2	0.139
	333653	AI051696		EST singleton (not in UniGene) with exon hit	0.139
		AA595289		EST singleton (not in UniGene) with exon hit	0.139
		AA013001	Hs.60563		0.139
45	329568		113.0000	CH.10_p2 gi 3962490	0.139
		AA253074	Hs.146261		0.139
	334320	70 2300. 7		CH22_FGENES.374_5	0.139
		Al916949	Hs.149748	ESTs; Weakly similar to weak similarity to collagens [C.elegans]	0.139
		AA864533		EST singleton (not in UniGene) with exon hit	0.139
50		AA984364	Hs.119064	ESTs	0.139
	*****	M99439	Hs.83958	transducin-like enhancer of split 4; homolog of Drosophila E(sp1)	0.139
		A1076204	Hs.135440		0.139
		AW072970		EST singleton (not in UniGene) with exon hit	0.139
	303273	AA316069		EST duster (not in UniGene) with exon hit	0.139
55	317484	AW274696	Hs.143921		0.139
	333239			CH22_FGENES.111_1	0.139
		Al184951		EST singleton (not in UniGene) with exon hit	0.139
	316813	AA826505	Hs.124517		0.139
60		AA281365	Hs.121640	ESTs; Weakly similar to KIAA0386 [H.sapiens]	0.139
60		AI700145		poly(A)-binding protein; cytoplasmic 1	0.139
		AW086142	Hs.159017		0.139
		AA335715	Hs.200299		0.139
		Al318342		EST singleton (not in UniGene) with exon hit	0.139
15	-	Al318327		EST cluster (not in UniGene)	0.139
65	327934		11- 40-400	CH.06_hs gij5868184	0.139
		AA670052	ms.195188	glyceraldehyde-3-phosphate dehydrogenase	0.139
	334756	A A 454 00T	He ones	CH22_FGENES.428_5	0.139
		AA451867	Hs.99255 He 150808	ESTs ESTs; Weakly similar to KIAA0644 protein [H.sapiens]	0.139 0.139
	301383	AI474722	113.130030	CO. 21 LIGUIN SHIMM IN UNIONAL PLANE IN LITTLE AND	U. 103

	312005	T78450	Hs.13941	ESTs	0.139
	338431			CH22_EM:AC005500.GENSCAN.351-4	0.14
		T90496	Hs.16757		0.14
_	333601			CH22_FGENES.213_4	0.14
5		AA278449	Hs.137429		0.14
	336911			CH22_FGENES.344-4	0.14
	338157			CH22_EM:AC005500.GENSCAN.209-5	0.14 0.14
	327845	745000	U- 00707	CH.05_hs gi 6531962	0.14
10		Z45662	Hs.90797	Homo sapiens clone 23620 mRNA sequence CH22_FGENES.428_12	0.14
10	334763 329384			CH.X_hs gi 5868869	0.14
		AF054663	•	EST cluster (not in UniGene) with exon hit	0.14
		AW452656	Hs.209824		0.14
	329916	AV1402000	110,200024	CH.16_p2 gi 6223624	0.14
15		N49826	Hs.18602	ESTs	0.14
	338129	1110020		CH22_EM:AC005500.GENSCAN.197-2	0.14
	325704			CH.14_hs gij5867028	0.14
	335656			CH22_FGENES.590_7	0.14
		W72366	Hs.40033		0.14
20	316807	AI018331	Hs.172444	ESTs; Highly similar to transcription regulator [M.musculus]	0.14
	310743	AW449754	Hs.158665	ESTs	0.14
	326941			CH.21_hs gi 6004446	0.14
	328809			CH.07_hs gi 5868327	0.14
	323855	Al653164	Hs.128665		0.14
25		AA564064		EST singleton (not in UniGene) with exon hit	0.14
	325666			CH.14_hs gi 6469822	0.14
	333747			CH22_FGENES.265_6	0.14
		AW015616	Hs.143321		0.141
20	332972	4.4007000		CH22_FGENES.51_5	0.141
30		AA825266	Un 100102	EST singleton (not in UniGene) with exon hit	0.141 0.141
		AW182805	U2' 193 193	ESTs; Weakly similar to Nod1 [H.sapiens]	0.141
	327296 336400			CH.01_hs gi 5867492 CH22_FGENES.823_15	0.141
		H26214	He 20733	ESTs; Weakly similar to !!!! ALU SUBFAMILY SX	0.141
35	321033	F1202 14	113.207.00	WARNING ENTRY	0.141
73	316522	AI475995	Hs.122910		0.141
	335715	71470000	110.122010	CH22_FGENES.599_15	0.141
	335959			CH22_FGENES.650_2	0.141
	333259			CH22_FGENES.118_7	0.141
40	337382			CH22_FGENES.744-8	0.141
	322346	AA227618	Hs.10882	HMG-box containing protein 1	0.141
	325378			CH.12_hs gi 5866920	0.141
	338500			CH22_EM:AC005500.GENSCAN.390-1	0.141
4 5	338460			CH22_EM:AC005500.GENSCAN.362-5	0.141
45		AW511138	Hs.256581		0.141
		AI539443	Hs.137447		0.141
	333624			CH22_FGENES.222_3	0.141
	329237			CH.X_hs gij5868729	0.141 0.141
50	330117			CH.19_p2 gi[6015201 CH22_EM:AC005500.GENSCAN.134-1	0.141
JU	338017			CH22_EM:AC005500.GENSCAN.134-1 CH22_EM:AC005500.GENSCAN.38-12	0.141
	337854 329984			CH.16_p2 gi 4646193	0.142
		AA622328	Hs.162762		0.142
		N40373	110.102704	EST cluster (not in UniGene) with exon hit	0.142
55	327823	1140070		CH.05_hs gij5867968	0.142
	326753			CH.20_hs qi 5867616	0.142
		AA904482	Hs.197775	ESTs	0.142
	334303			CH22_FGENES.373_6	0.142
42.2	326453			CH.19_hs gi 5867399	0.142
60		Al864581	Hs.215477		0.142
	308740	AI802711	Hs.210337	EST; Weakly similar to aldolase A [H.sapiens]	0.142
		H63959	Hs.142722		0.142
	338010			CH22_EM:AC005500.GENSCAN.128-8	0.142
	336326			CH22_FGENES.812_4	0.142
65		R44308	Hs.242302		0.142
		R55421		EST cluster (not in UniGene)	0.142
	325855	1104700 1741	704	CH.16_hs gl 5867067	0.142
	330425	HG1728-HT17	134	Non-Specific Cross Reacting Antigen (Gb:D90277), Alt. Splice Form 2	0.142
				rue opilos i utili e	U.142

	324583	AA425411		ESTs .	0.142
	326268			CH.17_hs gi 5867267	0.142
		AA460341		ESTs	0.142
5	338904			CH22_DJ32I10.GENSCAN.10-16	0.143 0.143
3	333096	AA446869	Hs.119316	CH22_FGENES.79_1	0.143
		Al248004	Hs.125187	_	0.143
		AW179174		ESTs	0.143
		Al204001		ribosomal protein L31	0.143
10		AA885317	Hs.190511		0.143
	337392			CH22_FGENES.747-3	0.143
	325543			CH.12_hs gi 6682452	0.143
		AA873085		EST singleton (not in UniGene) with exon hit phosphodiesterase l/nucleotide pyrophosphatase 2 (autotaxin)	0.143 0.143
15	332707	L35594	NS.174100	CH22 EM:AC005500,GENSCAN.59-10	0.143
13	337913	AA961061	Hs.131696		0.143
	335078	74201001	713.101000	CH22_FGENES.486_5	0.143
	338451			CH22_EM:AC005500.GENSCAN.359-39	0.143
	302777	AJ230640		EST cluster (not in UniGene) with exon hit	0.143
20		J03068		N-acylaminoacyl-peptide hydrolase	0.143
		H41411	Hs.33855		0.143
	328939	A1440474	U- 000007	CH.08_hs gi 6004481	0.143
	308015	Al440174	HS.228907	EST; Wealdy similar to GUANINE NUCLEOTIDE-BINDING PROTEIN BETA SUBUNIT-LIKE PROTEIN	
25				12.3 [H.sapiens]	0.143
23	328504			CH.07_hs gi 5868471	0.143
		AA402891	Hs.32951	solute carrier family 29 (nucleoside transporters); member 2	0.143
	335744			CH22_FGENES.601_15	0.143
		AF077208		EST cluster (not in UniGene)	0.143
30		AL042661		EST cluster (not in UniGene)	0.143
	318443	A1939323	Hs.15//14	ESTs; Weakly similar to NEURONAL ACETYLCHOLINE RECEPTOR PROTEIN; ALPHA-5 CHAIN PRECURSOR	
				[H.sapiens]	0.143
	336568			CH22_FGENES.843_7	0.143
35		H08815	Hs.159824		0.143
	327672			CH.04_hs gi 5867843	0.143
	335900			CH22_FGENES.635_8	0.144
	336044			CH22_FGENES.679_6	0.144
40	318845	Al815951		ESTs; Weakly similar to estrogen-responsive finger protein;	0.144
40	333483		efp [H.sapi	ensj CH22_FGENES.165_2	0.144
	333337			CH22_FGENES.139_6	0.144
		AA889197		EST singleton (not in UniGene) with exon hit	0.144
	335719			CH22_FGENES.599_22	0.144
45	325682			CH.14_hs gi 6138923	0.144
	327350			CH.01_hs gi 6249563	0.144
	339291			CH22_BA354112.GENSCAN.18-1	0.144 0.144
	326358			CH.18_hs gi 5867293	0.144
50	330316	Al499346	He 17/131	CH.08_p2 gi 6007576 ribosomal protein L6	0.144
50	338065	71433040	113.174101	CH22_EM:AC005500.GENSCAN.164-1	0.144
	339009			CH22_DA59H18.GENSCAN.18-7	0.144
	327776			CH.05_hs gi 5867964	0.145
	336664		•	CH22_FGENES.41-8	0.145
55		AF070619		EST cluster (not in UniGene)	0.145
		T70147	Hs.12024	ESTs EST singleton (not in UniGene) with exon hit	0.145 0.145
		AA062892	Hs.250178		0.145
	303818	Z45986	119:530110	CH.02_hs gi 6017023	0.145
60	335227			CH22_FGENES.513_13	0.145
00	339022			CH22_DA59H18.GENSCAN.22-1	0.145
		H55661	Hs.33026		
			TRAB [C.el	egans]	0.145
CF		AI697008	Hs.201811	Llare and an absence one 40: servid B00004	0.145 0.145
65		AA262760	ms.156015	Homo saplens chromosome 19; cosmid R29381 EST cluster (not in UniGene) with exon hit	0.145
	303252	AA156760		CH22_FGENES.757-2	0.145
		A1734009	•	EST cluster (not in UniGene)	0.145
	329333	3 1040		CH.X_hs gij5868806	0.145
				- ·	

	336857			CH22_FGENES.291-7	0.145
	332565	AA234896	Hs.25272	E1A binding protein p300	0.145
	318634	Al928098	Hs.156832	ESTs	0.145
	336318			CH22_FGENES.801_1	0.145
5		Al923551	Hs.170843		0.145
•	335346			CH22_FGENES.537_2	0.145
		T65416	Hs.12826	ESTs	0.145
	337607	100410	110.12020	CH22_C20H12.GENSCAN.17-3	0.146
		T84096	Hs.15284	ESTs	0.146
10			115.10204		0.146
10		T80698		EST cluster (not in UniGene) with exon hit	
		AA210878	11-050004	EST cluster (not in UniGene)	0.146
		Al695374	Hs.256231		0.146
		AA632554	Hs.163401		0.146
1.5		Al142359	Hs.155316		0.146
15		N70088	Hs.138467		0.146
	329276			CH.X_hs gi 5868762	0.146
	335887			CH22_FGENES.633_1	0.146
	338294			CH22_EM:AC005500.GENSCAN.297-1	0.146
	336993			CH22_FGENES.409-4	0.146
20	334135			CH22_FGENES.336_2	0.146
	326251			CH.17_hs gi 5867263	0.146
	337396			CH22_FGENES.749-1	0.146
	339167			CH22_DA59H18.GENSCAN.69-8	0.146
		AW135418	Hs.161210	. 	0.146
25	325313			CH.11_hs gi 5866865	0.146
		N66918	Hs.32205	ESTs	0.146
		AL043362		EST cluster (not in UniGene)	0.146
		AF062275		EST cluster (not in UniGene) with exon hit	0.146
		AA947909		EST singleton (not in UniGene) with exon hit	0.146
30	334399	70047505		CH22_FGENES.382_5	0.146
50	326472			CH.19_hs gij5867404	0.146
	333061			CH22_FGENES.75_4	0.146
	337072			CH22_FGENES.448-5	0.146
	334328			CH22_FGENES.375_5	0.146
35				CH.21_hs gi[6531965	0.146
<i>JJ</i>	327039			CH.12_hs gi[6552443	0.147
	325576	AIOTEONA	Hs.132660		0.147
		A1075804	115.152000	· · · · · · · · · · · · · · · · · · ·	0.147
		AA323758		EST duster (not in UniGene)	0.147
40	334501			CH22_FGENES.397_17	0.147
70	338238	A1744069		CH22_EM:AC005500.GENSCAN.264-4 EST circleton (not in UniCone) with even hit	0.147
	336567	Al744063		EST singleton (not in UniGene) with exon hit CH22_FGENES.843_6	0.147
					0.147
	335819			CH22_FGENES.619_2	0.147
45	336950	A14 40 477		CH22_FGENES.361-8	
43		Al148477	11- 400744	EST singleton (not in UniGene) with exon hit	0.147
		AW504854	Hs.126714	· ·	0.147
	335834			CH22_FGENES.621_1	0.147
	327870			CH.06_hs gij5868131	0.147
50		AA332011	HS.250138	protein phosphatase 2C; magnesium-dependent; catalytic subunit	
50	329412	44000000		CH.X_hs gi[6682553	0.147
		AA333068		EST cluster (not in UniGene)	0.147
		AA385315		EST duster (not in UniGene)	0.147
	327865			CH.06_hs gi 5868130	0.147
	333445			CH22_FGENES.154_2	0.147
55		AA021351	Hs.158497	KIAA0724 gene product	0.147
	336744			CH22_FGENES.118-9	0.147
		AA323414		EST duster (not in UniGene)	0.148
	320289	H07989		EST cluster (not in UniGene)	0.148
	305488	AA749000		EST singleton (not in UniGene) with exon hit	0.148
60	305592	AA780594	Hs.62954	ferritin; heavy polypeptide 1	0.148
	304094	H11295		EST singleton (not in UniGene) with exon hit	0.148
	325040	AW296368		EST cluster (not in UniGene)	0.148
	339034			CH22_DA59H18.GENSCAN.26-2	0.148
	334504			CH22_FGENES.398_2	0.148
65	334778			CH22_FGENES.431_2	0.148
		U77494	Hs.119687	RAN binding protein 8	0.148
		AW173759	Hs.203401		0.148
	325826			CH.15_hs gi[5867048	0.148
		T55182	Hs.152571	ESTs; Highly similar to IGF-II mRNA-binding protein 2 [H.sapiens]	
					-

	325785			CH.14_hs gij6381957	0.148
	333166			CH22_FGENES.91_8	0.148
	336548			CH22_FGENES.841_5	0.148
_	337552			CH22_C4G1.GENSCAN.1-4	0.148
5	331775	AA382742	Hs.97151	EST	0.148
	338936			CH22_DJ32I10.GENSCAN.19-6	0.148
	331869	AA428554	Hs.104894	ESTs; Weakly similar to fibronectin precursor [H.sapiens]	0.148
	332865			CH22_FGENES.28_5	0.148
40	328663			CH.07_hs gi 6004473	0.148
10	328436			CH.07_hs gi 5868417	0.148
		A1634864	Hs.250789	ESTs; Highly similar to similar to NEDD-4 [H.sapiens]	0.148
	336942		,	CH22_FGENES.354-2	0.148
		R53169	Hs.246091		0.149
15	333296			CH22_FGENES.132_3	0.149
15	333365		11- 050054	CH22_FGENES.142_2	0.149 0.149
		AW452392	Hs.252854		0.149
	337109	A14470000 .	Hs.190201	CH22_FGENES.489-2	0.149
		AW173300	NS. 190201		0.149
20	333454			CH22_FGENES.157_3 CH22_FGENES.432_9	0.149
20	334784			CH22_FGENES.118_3	0.149
	333255 337518			CH22_FGENES.814-7	0.149
		AA489268		EST cluster (not in UniGene)	0.149
		AA287567		EST cluster (not in UniGene)	0.149
25	328761	70201001		CH.07_hs gi[5868302	0.149
	328787			CH.07_hs gi[5868309	0.149
	335261			CH22_FGENES.520_2	0.149
		R16689	Hs.106004	_	0.149
	339263			CH22_BA354112.GENSCAN.10-1	0.149
30	337412			CH22_FGENES.756-6	0.149
	334414			CH22_FGENES.384_1	0.149
	332931			CH22_FGENES.38_5	0.149
		AW270980	Hs.106346	novel centrosomal protein RanBPM	0.149
0.5		AA669056		EST singleton (not in UniGene) with exon hit	0.149
35		AA470122	Hs.190261		0.149
	338414	****		CH22_EM:AC005500.GENSCAN.341-27	0.149 0.149
		AW247361		EST cluster (not in UniGene) with exon hit	0.149
	337509	41004440		CH22_FGENES.806-4	0.149
40		Al001149	Un 0/0116	EST singleton (not in UniGene) with exon hit chemokine (C motif) XC receptor 1	0.149
40	336536	L36149	NS.240110	CH22_FGENES.839_18	0.149
		T32458	Hs.14285		0.149
		Al767433	Hs.170013		0.149
	333595	707700	113.170010	CH22_FGENES.211_2	0.149
45	335975			CH22_FGENES.652_9	0.15
		AI003654		EST singleton (not in UniGene) with exon hit	0.15
	335025			CH22_FGENES.475_3	0.15
	328711			CH.07_hs gi 5868271	0.15
	328274			CH.07_hs gi 5868219	0.15
50	325505			CH.12_hs gi 6682451	0.15
	329641			CH.14_p2 gi 6468233	0.15
	304955	AA613504		EST singleton (not in UniGene) with exon hit	0.15
	339103			CH22_DA59H18.GENSCAN.44-10	0.15
	329636			CH.12_p2 gi 5302817	0.15
55		Al203293	Hs.157489		0.15
	326056	4.4700077		CH.17_hs gij5867184	0.15 0.15
		AA769074	Ho goos	EST cluster (not in UniGene) with exon hit	0.15
	303153	U09759	Hs.8325	mitogen-activated protein kinase 9	0.13

10

TABLE 13A shows the accession numbers for those primekeys lacking unigeneID's for Table 13. For each probeset we have listed the gene cluster number from which the oligonucleotides were designed. Gene clusters were compiled using sequences derived from Genbank ESTs and mRNAs. These sequences were clustered based on sequence similarity using Clustering and Alignment Tools (DoubleTwist, Oakland California). The Genbank accession numbers for sequences comprising each cluster are listed in the "Accession" column.

10	Pkey: CAT number: Accession:		Unique Eos probeset identifier number Gene cluster number Genbank accession numbers
15	Pkey	CAT number	Accession
		24275_1 1599424_1	AL137589 AA423949 BE222949 BE222694 Al199615 AW873116 Al277950 AW044290 AW630096 H61962 W01567 N75711
20		13653_22	BE259906 AA232518 AA013359 AL035788 AW160822 BE387134 BE002954 BE391839 AW161565 Al878841 BE616458 BE409981 BE387308 BE297436 BE315536 AA206924 R12012 AA214169 BE312812 BE387093 H11710 BE312009 BE260569 AA343566 AA219526 R34757 AA219749 BE336733 AA219751 AW411099 AA232408 BE018716 BE398089 AA206253 AA053487 AA114224 AV655868 AW732566 BE394087 AW732574 AA313442 BE336875 AA070548 BE259840
25			BE019828 AW732341 AA299916 BE019253 BE018238 BE387109 AA232304 BE255589 AW732585 AA181436 AA308777 AA075802 AW732521 AA314526 AA226747 BE409513 AA206168 BE388292 BE298782 BE387086 AA305310 AV652723 AA314918 BE615510 AW951763 BE398104 BE385195 BE407165 BE391336 BE390187 BE389189 BE540650 BE249884 BE385985 BE274245 BE391124 BE260080 AA182600 BE512821 BE390090 BE279398 BE279589 BE263454 BE515194 BE293569 BE272531 BE388814 BE384659 BE271685 BE561043 BE278449 BE302572 AW239076 AI750583 AA376179 AA112632 BE266324 BE266614 R13105 AA132286 BE296305 Al220355 AA205606 AA219527 AA219519 AW804310
30	322303	622937_1 704603_1	AA083286 BE171208 T19693 AA338328 BE185868 AA903024 T92162 AA330119 BE410404 BE314668 AW576245 BE207878 AW299993 Al199558 Al285442 AW299994 AW394242 AW394184 Al357412 Al870708 Al590539 W07459 AW068287 AA310079 BE336702 AA356318 AA306059 AA346785 AW402633 AA311210 AW402909 N76879 AW402913
35	322394	27492_1	AW401920 AA321636 AA354474 C17297 C16938 AA311774 M29871 NM_002872 Z82188 AW405674 H94176 R89281 AA214723 AI014482 AW949347 T27749 AW804226 AW796964 AW404581 AF077208 NM_014029 W68830 W79652 AA353375 AW575218 AA552192 AA521232 AA702695 AA033975 AW407827 AA829948 N94402 AW628604 AI523308 N57605 AA641662 H42477 N52784 AI753478 AA768493 AA845729 W47391 N55270 AI090117 R89282 BE206172 AA076650 AA595650 AI218931 BE049397 AI433110 W74114 H94277 AI358627 AI085221 AI862818 AA835967 AW103905
40	201750	44275_1	A1640644 AA835507 AA856887 AA694392 AW337542 A1524410 BE045500 A1440060 A1358801 AW028238 AW205248 A1718264 R48618 AA357358 A1695002 AA897549 AW081065 A1433360 A1810783 A1620963 Z82188 AA360224 U29112 A1656540 A1364875 A1656246 A1990940
		155498_1	AA169345 AI762857 AI949997 AI809601 AI681948 AI221079 AW167404 AI347614 AI611090 AI023472 AI347683 AI027467 AW591788 AI380665 AA835735 AA836654 AI244028 AW193159 AI500112 AI918722 AI738693 AI702308 AA805365 AI766842
45	321921	38937_1 34680_1 21620_1	T59538 T59589 T59598 T59542 AF147374 AF070619 R20302 T80358 AJ223366 BE305086 AW820106 AA621983 BE305208 AI738475 AI380189 AW590847 AI127232 AA622706 AI380858 AA621975 AI587036 AA665743 AW204003 AI692234 AI002242 AI692219 AW137282 AW268783 AW295910 AI308015
50			AW301462 Al318288 Al318575 Al318117 Al345591 Al249650 Al246934 Al246864 Al246971 AW268311 Al249654 BE041907 AW732776
		265316_1 14694_7	N72324 N52825 W19526 BE143464 AA376060 M83667 NM_005195 S63168 M83667 AW068039 AW630649 Al338577 Al018125 Al269878 AW242440 Al887823 Al342581 BE222416 Al582847 Al651011 Al660815 Al699574 BE550201 Al926996 AW665855 Al827752 Al761857 BE328168 BE222451 Al762201 AW000929 AW007207 BE042962 BE551843 BE465373 Al279179 Al949945 BE551862 AW051667
55			BE328076 BE222296 AW007229 AW0772332 AI279801 AI334526 AI631938 AI770103 BE041412 AI417900 AI692655 AI869943 AW270119 AI431739 AI703347 AW770568 AW025473 AI701497 AI128026 BE328147 AW203980 BE046793 AW087704 AI674597 AI650732 AI813691 AI472092 AI695224 AI241217 AW207746 AI206840 AI271362 AI631788 AI911883 AI914619 AI380585 AI767501 AI823759 AI564116 AI190991 AI377369 AI814122 AI221623 AI354793 AI081988 AI391740 AI337435 BE467366 AI824347 AI565325 AI280038 AI640455 AI819744 BE467803 BE327524 AI149402 AI313187 BE219684
60			AW611948 AW665821 Al091260 AW044492 BE220366 AW025381 AW183264 Al694865 Al498474 Al129780 Al202028 Al566792 BE220659 Al928040 Al830696 Al493021 AW612488 Al913152 BE042965 Al631837 Al693873 Al498925 Al768668 Al401544 BE327023 Al693383 Al769874 Al744003 AW082273 Al686501 Al798177 Al985196 Al090033 Al432342 Al689918 Al638308 BE468080 BE219588 Al912119 BE219787 AW005392 BE326564 Al589039 Al860187 Al758143 Al338168 Al702936 BE221985 Al498727 Al918196 Al279735 AW771497 Al860133 AW237834 AW661759 AW028111 BE503416
65 '			Al360180 AW611715 Al871777 BE045447 BE326444 Al266547 Al800237 Al823315 Al478368 Al264281 Al675841 Al690041

		A1498018 AI554124 AI239893 AI864054 AI280099 AI192815 AI620465 AI080201 AW002057 BE500986 AI341131 AI818991 AI566137 AI123403 BE219192 AW183844 AI499842 AW137971 AW138720 AW015526 AW138160 AW243163 AW138705 AW139927 AW140006 AW138810 AW137450 AW206970 AW135419 AW205974 AA043494 BE465106 AW139955 AI741112 BE326942 AA043506 AI079957 AI942432 AI392902 AI097047 AI470599 AA514553 AA984008 N47949 AI654114 AA884832
5	· .	A1796752 A1765290 Al301155 AW470358 BE222764 Al823569 Al651188 Al692695 Al476643 BE504307 Al767573 BE219719 Al936249 AW467075 Al913633 BE221966 Al091025 AA969215 A1799810 AA931170 BE048559 Al809606 Al138614 A1739456 Al674605 AW772068 Al089286 Al625787 Al263418 AW008638 Al928389 AW628997 Al470010 Al914168 Al760003 Al203050 Al334069 Al694788 BE045337 Al948659 Al912982 Al867131 Al192102 Al767583 Al347518 Al566005 Al625884 Al215888 Al633904 AW182265 AW614357 Al128030 Al343685 Al914283 Al985003 Al823578 Al493053 Al380285 Al633895
10		Al267880 Al538162 Al991552 BE219479 BE219296 Al302178 AW779296 Al913805 Al631644 Al566772 Al985498 Al942289 Al935659 Al339092 Al247432 Al686472 Al766886 Al017228 Al333272 AW301668 Al972218 AW082027 Al632974 Al474761 Al766127 AW236578 AW000966 Al870734 Al222399 Al871249 Al703448 BE464210 Al768037 Al871585 Al767871 Al738757 Al220732 Al681633 Al768783 Al684463 Al307339 Al263203 AW665264 BE463969 Al768786 Al439118 Al127913 BE218324
•		AI672342 BE220052 AI796163 AI221662 AW197672 AW025300 AI769681 AW612448 BE219757 AW072420 AI669980
15		Al830418 AW204353 AA047011 AA913868 AI739146 Al669954 AW470507 AW614835 AW302151 AW772372 AI762427 AW339902 AW303370 BE464775 AW299818 AW236072 AW195060 AW274737 AW263062 AW183846 Al868894 AW300493 AW172509 AW516876 AW593773 AW299474 AW303546 Al817323 Al823624 Al694005 Al934589 Al343479 Al861825 Al962726 AI765845 AW080318 Al640227 AI763042 AI768903 AW235386 AA738489 AW341293 AA588585 BE221732 Al914179 AW611669 AI572789 AW194735 AW236122 AW236007 AW612789 AW197501 AW195046 AI797145 Al864423
20	• •	A1914179 AW011069 A1572789 AW194733 AW230122 AW23007 AW012769 AW197301 AW19500 A127170 A1458934 A1342848 A1693227 A1912642 A1689993 AA9322572 AA740269 AW470392 AW086020 A1221701 T69326 T70461 A1765579 A1338263 A1431721 A1394249 A1186462 A1823571 A1533665 A1497954 A1761057 A1678228 A1640302 A1948742 AA594626 AA883155 A1972682 A1804774 A1300407 A1433524 AA897341 A1401175 A1291071 AA021213 A1126509 A1948955
		AI218835 AA903938 AA502610 AI498320 AA584267 AA935285 AI476253 AA489658 AA975053 AA715326 AA557139 AA126417 AA971455 AA557319 AI499738 AA911438 AI913637 AA494506 N90793 AI990724 AA131667 AA128164
25		AA046840 Al262557 AA131729 AA594926 T59467 AA436907 AA044630 Al589177 Al279237 Al880498 Al431822 AA708934
		AW612558 Al634069 W03610 Al192272 BE550862 Al400879 AA708507 Al128003 Al375308 Al271423 Al199552 AA125977
		Al366498 AA458662 Al694382 AA044627 Al636263 Al796270 T90146 AW014724 Al870812 Al948781 AA369965 Al094721
		AW271817 Al262898 Al244680 T69252 Al934148 AA046357 W19109 AA028157 AW021924 AA253491 Al189397 Al934388
30		D58282 W21323 W24288 AI682972 AA293683 AA284566 AV659511 AA434184 H87089 AA040038 N57464 AA343709 AW805815 R89837
30	301119 33384_1	BE621320 BE266806 BE276582 AW516729 AF142579 AW451687 AK000069 AA325236 BE168997 W73105 AA715365
	301119 30304_1	BE278873 AA808894 AA386371 AW517942 AW750993 BE140314 BE392384 BE621757 AA318192 BE548173 AW152607
		AW166898 AA352215 AW841506 T59802 AF147378 AA335719 AW956069 T59668 AA826362 Al961329 Al290469
		AW197375 AI805651 AA160748 AA581089 AI968889 AA581100 AA501478 AI621069 AA488534 AA503715 AA658457
35		Al144504 BE387827 AA159880
	324019 262792_1	AW177009 AJ381610
	323437 189513_1	AA287567 AA252404 AW967735 AA287568 AA761222 AA865644 AA831245
	307845 19804_10	BE514807 R43224 Al363450 AA450226 AF030942 AA385315 Al627453 Al050695 Al348281
40	324126 272259_1 309101 7570_1	AI340462 AI583268 AA079086 AI950777 AI301866 AI925108 AW876954 AW877000 AA525418 AA888549 AI934220
70	209101 7570_1	AW380220 AA804858 AI927576 T61151 AW384053 BE391691 AA533856 AA248400 T48202 N57156 R68346 R26020
		ALD50332 W30806 H61369 AAD92592 AA230324 BE271217 AW372903 T48772 AA358002 AA094302 AA559856 AW373308
		AW373315 AW373297 AW373311 AW373314 AW373309 AW877055 AW770140 AW379805 Al581609 AW364144 AA078921
		AA715432 AA654210 Al004899 AA602209 W47464 AA506588 R26822 AU076528 AI535743 AI535704 AI535681
45	315703 119175_1	AA402307 D60405 D61237 D59891 AW964877 AA325215 Al459739 N36070 N25658 AA083684 AW293368 AI761958 AI741205 Al693175 AW873603 AI143269 AI187124 N25199 H19323 Al650842 AW316825 AA083842 AA935650 AW298404
		A1741205 A1693175 AW873603 A1143269 A1167124 N25199 H19323 A1630642 AW316625 A4063642 A4953660 AW436444 A1672001 A1648568 R17676 R41625 A1123237 R17677 A1206866 F36920 A1654713 F34084 AA618029 A1915139 AW275194
		AW514577 D80420 AW149850 Z40953 AI867861 AA927547 AA974344 AI825793 AI635565 AI652157 BE504748 AW295759
		F16800 AW839796 F01781 AA909730 AA984010
50	301373 368214 1	AA595235 AW973839 T03040
	323665 54093_1	AW248307 AA313452 AW951927 AA355961 BE566080
	323665 54093_1 323676 220254_1	AI702835 AI758919 AI685405 AI952108 AI299207 AI400767 AW105389 AI952710 AA845312 AI784118 AI537315
	323665 54093_1 323676 220254_1 302086 23306_1	AI702835 AI758919 AI685405 AI952108 AI299207 AI400767 AW105389 AI952710 AA845312 AI784118 AI537315 X57138 NM_003514 Z98744 BE253911 BE256314 AI095013 AI138475
55	323665 54093_1 323676 220254_1 302086 23306_1 323731 226193_1	AI702835 AI758919 AI685405 AI952108 AI299207 AI400767 AW105389 AI952710 AA845312 AI784118 AI537315 X57138 NM_003514 Z98744 BE253911 BE256314 AI095013 AI138475 AA323414 AW664013 AI809377 AI276041 AW296883 AI798340
55	323665 54093_1 323676 220254_1 302086 23306_1 323731 226193_1 323791 232336_1	AI702835 AI758919 AI685405 AI952108 AI299207 AI400767 AW105389 AI952710 AA845312 AI784118 AI537315 X57138 NM_003514 Z98744 BE253911 BE256314 AI095013 AI138475 AA323414 AW664013 AI809377 AI276041 AW296883 AI798340 AA333068 AA331863 AA331838 AW962531 AA331442
55	323665 54093_1 323676 220254_1 302086 23306_1 323731 226193_1	AI702835 AI758919 AI685405 AI952108 AI299207 AI400767 AW105389 AI952710 AA845312 AI784118 AI537315 X57138 NM_003514 Z98744 BE253911 BE256314 AI095013 AI138475 AA323414 AW664013 AI809377 AI276041 AW296883 AI798340 AA333068 AA331863 AA331838 AW962531 AA331442 AW296368 AA247632 AK002030 R15304 T08775 AW975664 AI186801 AA730688 AW190918 AI141176 AW513211
55	323665 54093_1 323676 220254_1 302086 23306_1 323731 226193_1 323791 232336_1 325040 23854_1	AI702835 AI758919 AI685405 AI952108 AI299207 AI400767 AW105389 AI952710 AA845312 AI784118 AI537315 X57138 NM_003514 Z98744 BE253911 BE256314 AI095013 AI138475 AA323414 AW664013 AI809377 AI276041 AW296883 AI798340 AA333068 AA331863 AA331838 AW962531 AA331442 AW296368 AA247632 AK002030 R15304 T08775 AW975664 AI186801 AA730688 AW190918 AI141176 AW513211 AI275071 AA988601 BE042933 BE045713 AW087176
55	323665 54093_1 323676 220254_1 302086 23306_1 323731 226193_1 323791 232336_1 325040 23854_1 324430 312113_1	AI702835 AI758919 AI685405 AI952108 AI299207 AI400767 AW105389 AI952710 AA845312 AI784118 AI537315 X57138 NM_003514 Z98744 BE253911 BE256314 AI095013 AI138475 AA323414 AW664013 AI809377 AI276041 AW296883 AI798340 AA333068 AA331863 AA331838 AW962531 AA331442 AW296368 AA247632 AK002030 R15304 T08775 AW975664 AI186801 AA730688 AW190918 AI141176 AW513211
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20	306051 19085_3	F08147 AW408359 AW949429 R23785 AW247442 AA305512 T29095 AA905130 BE246361 BE244981 AA220199 BE504058 X80878 AA533727 AA608601 AW005964 Al811627 Al367037 Al277985 Al493719 Al277848 AA854982 AW247298 Al216345 Al041295 Al887378 AA781241 Al674270 AW628959 Al383083 BE504391 AA729421 AA552188 AA373387 AW880360
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		c11_hs	414/040040
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	304475		AA428879
	304522		AA465405
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	306004		AA889992
	306008		AA894390
	306013		AA896990
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	306249		AÅ933840
	306286		AA936892
25	306295		AA937331
25	306317		AA947909
	306347		AA961144 AA962086
	306365 306398		AA970548
		entrez D28383	D28383 •
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		•	Al304330 Al339217 AW193588 AW438688 Al818970 AW316799 AA906527 AA777570 N47673 Al336428 AW945133
			Al038606 R29692 AW194197 Al304748 H12639 AA053178 AA493213 AA676958 AA113154 Al313469 Al368239 R93183
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35	330535	1374 -8	U11872
		10404 2	U24488 NM_007116
			

TABLE 13B shows the genomic positioning for those primekeys lacking unigene ID's and accession numbers in Table 13. For each predicted exon, we have listed the genomic sequence source used for prediction. Nucleotide locations of each predicted exon are also listed.

Sequence source. The 7 digit numbers in this column are Genbank Identifier (GI) numbers

Unique number corresponding to an Eos probeset

5

Pkey:

Ref:

Pkey	Ref	Strand	Nt position ·	
	1101			
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3327	92 Dunham, I. et.al.	Plus	73381-73768	·
3328	10 Dunham, I. et.al.	Plus	304296-304384	
3329	44 Dunham, I. et.al.	Plus	2414825-2414932	
	72 Dunham, I. et.al.		2572152-2572236	
3331	33 Dunham, I. et.al.	Plus	3360058-3360195	•
3331	54 Dunham, I. et.al.	Plus	3615887-3616019	
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3332	27 Dunham, I. et.al.	Plus	3992866-3992968	
3332	30 Dunham, I. et.al.	Plus	3995507-3996507	
3332	98 Dunham, I. et.al.	Plus	4581537-4581947	
3333	04 Dunham, I. et.al.	Plus	4629943-4630242	
3333	05 Dunham, I. et.al.	Plus	4630388-4630645	
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	91 Dunham, I. et.al.		4916697-4916780	
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	97 Dunham, I. et.al.		4922466-4922635	
	03 Dunham, I. et.al.		4925140-4925256	
	13 Dunham, I. et al.		4943824-4943974	
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	·		6595146-6595244	
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r20		ERILLA	PM IN / 41-M316003	

6816731-6816993

6822087-6822406

6831369-6831445 6835282-6835474

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333656 Dunham, I. et al. Plus

333657 Dunham, I. et.al. Plus 333658 Dunham, I. et.al. Plus

65

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	336548	Dunham, I. et al.	Plus	34353881-34354826

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                                        34331316-34331183
        336542 Dunham, I. et.al. Minus
                                        34375244-34374907
        336556 Dunham, I. et.al. Minus
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                                        34375443-34375341
        336557 Dunham, I. et.al.
                                Minus
        336558 Dunham, I. et.al. Minus
                                        34375825-34375698
                                        34376430-34376261
                                Minus
        336559 Dunham, I. et.al.
                                        34376814-34376596
        336560 Dunham, I. et.al.
                                Minus
        336561 Dunham, I. et.al.
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                                        34377168-34376928
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                                        7627912-7627757
        336597 Dunham, I. et.al. Minus
        336601 Dunham, I. et.al.
                                Minus
                                         13265853-13265654
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                                         1304281-1304212
                                Minus
                                        1351268-1351168
        336645 Dunham, I. et.al.
                                        2158060-2157993
        336662 Dunham, I. et.al.
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        336664 Dunham, I. et.al.
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                                         1993558-1993481
        336676 Dunham, I. et.al.
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                                Minus
                                        2158060-2157993
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        336684 Dunham, I. et.al.
        336686 Dunham, I. et.al.
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                                        2160698-2160486
                                        3094026-3093871
        336714 Dunham, I. et.al. Minus
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        336719 Dunham, I. et.al.
        336736 Dunham, I. et.al.
                                Minus
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        336744 Dunham, I. et.al. Minus
                                        5419973-5419873
                                Minus
        336786 Dunham, I. et.al.
        336793 Dunham, I. et.al.
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                                        8201756-8201561
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                                Minus
                                        8396673-8396425
                                Minus
        336863 Dunham, I. et al.
        336933 Dunham, I. et al.
                                Minus
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         336942 Dunham, I. et.al.
                                Minus
                                         12027537-12027455
                                Minus
                                         13267243-13267172
        336960 Dunham, I. et.al.
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        336969 Dunham, I. et.al.
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                                         13725722-13725643
                                Minus
                                         13732308-13732221
        336971 Dunham, I. et.al.
                                Minus
                                         15523541-15523422
        337003 Dunham, I. et.al.
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        337011 Dunham, I. et.al.
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        337070 Dunham, I. et.al.
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                                         19657011-19656881
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                                         22649450-22649388
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                                Minus
                                         24594969-24594874
        337193 Dunham, I. et.al.
                                        27659956-27659876
         337256 Dunham, I. et.al.
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                                        28491414-28491094
        337284 Dunham, I. et.al.
                                Minus
                                        28846334-28845873
         337293 Dunham, I. et.al.
                                Minus
         337316 Dunham, Letal. Minus
                                         29657129-29656997
        337326 Dunham, I. et.al. Minus
                                        30017199-30017069
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	337382	Dunham Latel	Minus	31233666-31233579
	337392	Dunham, I. et.al. Dunham, I. et.al.	Minus	31442311-31442229
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	337406	Dunham, Let.al.	Minus	
5	337412	Dunham, I. et.al.	Minus	31916487-31916312
3	337419	Dunham, I. et.al.	Minus	32021496-32021170
	337436	Dunham, I. et.al.	Minus	32257869-32257739
	337455	Dunham, I. et.al.	Minus	32434517-32434425
	337509	Dunham, I. et.al.	Minus	33414613-33414498
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10	337529	Dunham, I. et al.	Minus	34043668-34043546
	337533	Dunham, I. et al.	Minus	34193388-34193261
	337539	Dunham, I. et.al.	Minus	34254490-34254322
	337551	Dunham, I. et.al.	Minus	34524446-34524362
	337553	Dunham, I. et.al.	Minus	24230-24160
15	337591	Dunham, I. et.al.	Minus	1006414-1006184
	337592	Dunham, I. et.al.	Minus	1007791-1007634
	337593	Dunham, I. et.al.	Minus	1009460-1009291
	337607	Dunham, I. et.al.	Minus	1355719-1355637
	337612	Dunham, I. et.al.	Minus	1570235-1570142
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	337824	Dunham, I. et.al.	Minus	4559540-4559266
	337825	Dunham, I. et.al.	Minus	4567155-4567005
	337850	Dunham, I. et.al.	Minus	5077143-5076943
	337854	Dunham, I. et.al.	Minus	5153435-5153272
25	337913	Dunham, I. et.al.	Minus	6149843-6149786
	337915	Dunham, I. et.al.	Minus	5922748-5922690
	337968	Dunham, I. et.al.	Minus	7095797-7095680
	338010	Dunham, I. et.al.	Minus	7754282-7754184
	338012	Dunham, I. et.al.	Minus	7761421-7761351
30	338017	Dunham, i. et.al.	Minus	7864521-7864401
	338065	Dunham, I. et.al.	Minus	7235048-7234950
	338094	Dunham, I. et.al.	Minus	9595602-9595440
	338129	Dunham, I. et.al.	Minus	10915338-10915237
	338132	Dunham, I. et.al.	Minus	10989617-10989530
35	338150	Dunham, I. et.al.	Minus	11478551-11478355
	338157	Dunham, I. et al.	Minus	11731444-11731375
	338195	Dunham, I. et.al.	Minus	13484103-13483972
	338255	Dunham, I. et.al.	Minus	15242294-15242231
	338276	Dunham, I. et.al.	Minus	16109555-16109398
40	338431	Dunham, I. et.al.	Minus	19747608-19747496
	338448	Dunham, I. et.al.	Minus	20151152-20151054
	338451	Dunham, I. et.al.	Minus	20174286-20174193
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	338684	Dunham, I. et al.	Minus	24827522-24827428
	338689	Dunham, I. et.al.	Minus	24893073-24892972
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50	338842	Dunham, I. et.al.	Minus	27824238-27824079
	338893	Dunham, I. et.al.	Minus	28491807-28491631
	338904	Dunham, I. et.al.	Minus	28766345-28766253
	338935	Dunham, I. et.al.	Minus	29071537-29071461
	339022	Dunham, I. et.al.	Minus	30523414-30523289
55	339034	Dunham, I. et.al.	Minus	30621603-30621422
	339190	Dunham, I. et.al.	Minus	32403103-32402985
	339212	Dunham, I. et.al.	Minus	32494335-32494210
	339213	Dunham, I. et.al.	Minus	32496590-32496440
	339216	Qunham, I. et.al.	Minus	32504250-32504109
60	339233	Dunham, I. et.al.	Minus	32751331-32751238
	339258	Dunham, I. et.al.	Minus	32934756-32934615
•	339262	Dunham, I. et.al.	Minus	32971258-32971090
	339263		Minus	32974634-32974452
	339265		Minus	32975943-32975806
65	339338		Minus	33468728-33468606
	339396			34017306-34017205
•	339400			34045024-34044940
	339425			34407911-34407798
	325207		Plus	140049-140170
		-		

	329568	3962490	Pius	36331-36750
		3983513	Minus	53197-53269
		5866865	Minus	27385-28192
5		5866875	Plus	75189-75264
3	325317	5866878	Minus	156551-156649 10867-10955
	325257 329632	5866895 6729060	Plus Plus	192813-193017
	325371	5866920	Minus	1035422-1035536
	325375	5866920	Minus	1165503-1165810
10	325378	5866920	Minus	1187981-1188167
	325469	6017034	Plus	286823-286991
	325470	6017034	Plus	287578-287663
	325576	6552443	Minus	137769-137894
15	325505	6682451	Minus Plus	240852-240946 151873-152057
13	325543 329635	6682452 5302817	Minus	62522-62622
	329636	5302817	Minus	64969-65078
	325593	5866992	Minus	469726-469860
	325675	5867014	Plus ·	955517-955711
20	325704	5867028	Plus	156198-156387
	325682	6138923	Plus	370618-370763
	325785	6381957	Plus	61849-62003
	325666	6469822	Plus	16769-16857 120278-120559
25	325818 329777	6682490 6002090	Minus Minus	191389-191479
23	329768	6015501	Plus	118315-118422
	329759	6048280	Minus	37647-37730
	329731	6065783	Plus	158772-158900
	329687	6117856	Minus	22165-22288
30	329676	6272128	Minus	142207-142359
	329667	6272129	Plus	101355-101745
	329669	6272129	Plus	131223-131291 131351-131495
	329670 329641	6272129 6468233	Plus Minus	105995-106107
35	329791	6469354	Minus	131982-132089
	325826	5867048	Minus	46361-46458
	325829	5867052	Plus	232674-233060
	329888	6067149	Minus	37227-37473
40	329893	6525313	Minus	166123-166791
40	329899 325988	6563505 5867064	Minus Plus	111058-111783 17349-17606
	325855	5867067	Plus	276141-276251
	325999	5867073	Plus	149115-149192
	326001	5867073	Pius	155223-155348
45	325886	5867087	Plus	194694-194915
	325882	5867087	Minus	8178-8347
	325905	5867104	Plus	78779-78876
	325922	5867122 5867132	Minus Minus	329063-329134 152633-152902
50	325937 325960	5867147	Minus	162506-162635
30	325961	5867147	Minus	165106-165209
	325838	6552452	Plus	171451-171532
	325839	6552452	Plus	181964-182037
بر ب	325840	6552452	Plus	184380-184547
55	325844	6552453	Minus	14188-14332
	325870	6682492 4646193	Plus Minus	228209-228297 139780-139890
	329984 329976	4878063	Minus	62584-62691
	329935	6165200	Minus	69059-69127
60	329916	6223624	Plus	36396-37195
•	330021	6671889	Plus	120938-121032
	330024	6671908	Minus	1005-1270
	330028	6671908	Minus	30015-30144
CE	326033		Plus	37261-37333
65	326036	5867178 5867184	Minus	120215-120273
	326056 326116		Minus Plus	181553-181690 45548-45604
	326122	5867194	Plus	144397-144683
	326138	5867203	Minus	179374-179436
			_	

	326145	5867204	Minus	52599-52814
	326180	5867211	Minus	182758-183222
	326201	5867216	Minus	166168-166959
5	326207 326226	5867222 5867230	Pius Pius	48139-48219 52644-52705
3	326233	5867232	Pius	124788-124863
	326238	5867260	Plus	64282-64338
	326241	5867260	Minus	181648-181916
10	326243	5867261	Plus	123838-123978
10	326251	5867263	Minus	82716-82822 122114-122765
	326268 326124	5867267 5916395	Plus Plus	407102-407560
	326339	6056311	Minus	164637-165251
	330049	4567182	Minus	314662-315210
15	326358		Plus	9122-9195
	326365	5867297	Minus	96630-96764
	326379 326382	5867327 5867327	Plus Minus	32299-32402 50420-50503
	326390		Minus	108814-110592
20	326424	5867369	Minus	168329-168409
	326453		Plus	86222-86423
	326472	5867404	Plus	293739-293940
	326492 326533	5867422 5867441	Plus Minus	120768-120991 532153-532280
25	330117	6015201	Minus	7340-7680
	330115	6015202	Plus	11403-11677
	330116	6015202	Plus	12109-12418
	330095	6015278	Plus	15343-15814
30	330096 326644	6015278 5867559	Plus Plus	49370-49458 42684-42819
50	326713		Plus	121511-121798
	326745	5867611	Plus	127130-127318
	326752	5867615	Minus	1214-1562
25	326753		Plus	12454-12511
35	326598	5867634 6552455	Plus Plus	68955-69014 142311-142441
	326667 326855	6552460	Minus	111390-111463
	326812	6682504	Plus	189811-189941
	327005	5867664	Plus	610847-610907
40	327008	5867664	Plus	928737-928811
	326896	5867680 5867684	Minus Minus	12032-12122 9280-9606
	326904 326951	6004446	Plus	193812-193998
	326941	6004446	Pius	62018-62896
45	326943		Minus	89242-89427
	326928		Minus	291007-291219
	326958	•	Minus Minus	42952-43082 43159-43301
	326959 327039	6531965	Plus	694486-694998
50	327127	6682520	Plus	41925-42083
	330158	6580367	Plus	81966-82456
	327204	5867447	Plus	165135-165239
	327208		Plus	180805-180864 82400-82615
55	327266 327277		Minus Minus	165616-165715
33	327289		Plus	49296-49536
	327296		Plus	7627-8166
	327237		Minus	59702-59813
<i>6</i> 0	327145		Minus	40482-40551 141448-141609
60	327333 327335		Minus Minus	142979-143124
	327343		Minus	12288-12395
	327350		Minus	41890-41985
<i></i>	327358		Minus	3802-3950
65	327360		Minus	6255-6422
	327409 327424		Minus Plus	52949-53011 160442-160598
	327424		Plus	1320-1403
	327470		Plus	150910-150973

	327460	6004455	Plus	175245-175343
	327498	6017023	Minus	42178-42283
	327509	6117815	Minus	54882-55053
_	327510	6117815	Minus	56824-56944
5	327512	6117815	Plus	176256-176325
'	327535	6525279	Plus	19105-19175
	330163	6042042	Minus	20321-20385
	330171	6648220	Plus	110889-111575
	327579	5867824	Minus	37229-38335
10	327672	5867843	Minus	69649-69740
	327629	5867872	Plus	49692-49811
	327640	5867890	Plus	9448-9566
	327649	5867899	Plus	205871-205927
	327612	6525283	Plus	2747-2924
15	327718	6525284	Plus	86123-86186
	327801	5867924	Plus	23239-23348
	327762	5867961	Minus	50303-50439
	327763	5867961	Plus	229347-229476
20	327776	5867964	Minus	164308-164486
20	327822	5867968	Minus	168886-169633
	327823	5867968	Minus	170359-170433
	327807	5867968	Plus	33745-33811
	327845	6531962	Plus	193402-193549 3719-3787
25	330228	6013527 6165182	Minus Plus	36103-36243
25	330190 328122	5868031	Plus	158474-158656
	328132	5868038	Minus	126737-126839
	328159	5868065	Minus	52957-53162
	328168	5868071	Plus	60321-60479
30	328175	5868073	Plus	208-271
	328217	5868096	Minus	3742-4362
	327865	5868130	Plus	61503-62205
	327866	5868131	Minus	2893-3046
	327870	5868131	Plus	53558-53757
35	327879	5868142	Minus	77722-77793
	327902	5868158	Minus	133339-133467
	327918	5868165	Plus	547530-547591
	327934	5868184	Plus	41830-42036 46497-46682
40	327959 327976	5868210 5868212	Minus Minus	349301-349409
40	328020	5902482	Minus	556386-556652
	328042	5902482	Minus	1985085-1986626
	328008	5902482	Plus	296663-297151
	330301	2905862	Minus	4420-5781
45	330299	2905881	Minus	1020-1382
	328274	5868219	Minus	31244-31439
	328595	5868224	Plus	148738-148967
	328591	5868227	Minus	237647-237726
~ 0	328668	5868254	Minus	10888-10984
50	328677	5868256	Minus	58708-58950
	328687		Plus	624479-624585
	328706		Plus	165501-165614
	328711	5868271	Minus	97797-97990
55	328730		Plus Plus	8068-8214 37437-37550
<i>JJ</i>	328732 328734		Plus	50559-50747
	328752		Minus	114911-115087
	328755		Minus	145959-146446
	328761		Minus	239308-239412
60	328775		Plus	12845-12920
.	328784		Minus	74523-74604
	328787		Plus	135772-135963
	328809		Plus	91792-91849
	328829		Plus	36309-36630
65	328280	5868352	Plus	160563-160631
	328311		Minus	170560-170826
	328318		Plus	414945-415620
	328323		Minus	1080089-1080235
	328348	5868383	Minus	260272-260379

	328377	5868390	Plus	16947-17023
	328436	5868417	Plus	203760-203904
	328504	5868471	Plus	47064-47217
	328506	5868471	Plus	60716-60830
5	328522	5868477	Plus	1972307-1972452
	328525	5868482	Plus	12387-14313
	328541	5868486	Plus	130956-131050
	328662	6004473	Plus	1184773-1184855
	328663	6004473	Plus	1185279-1186634
10	328803	6004475	Minus	291716-291948
	328304	6004478	Minus	3884-3952
	328927	5868500	Minus	428829-428893
	328936	5868500	Minus	1352202-1352259
	328939	6004481	Minus	131139-131320
15	328941	6456765	Minus	9817-9885
	328948	6456765	Plus	28227-28413
	328968	6456775	Plus	117442-118283
	330316	6007576	Minus	119761-119931
	330350	3056622	Minus	26413-26820
20	330351	3056622	Minus	27522-27614
	330348	4544475	Minus	19855-19962
	329034	*****	Minus	32819-32939
	329046	5868569	Plus	18971-19030
	329053	5868574	Pius	426453-426541
25	329186	5868711	Minus	13108-13225
	329237	5868729	Plus	133238-133339
	329276	5868762	Minus	222629-222709
	329333	5868806	Plus	392666-392746
00	329376	5868859	Plus	52356-52694
30 ·	329384	5868869	Minus	116524-116662
	329140	6017060	Plus	290842-290905
	329317		Plus	614823-615209
	329319	6381976	Plus	721390-721470
25	329129	6588026	Plus	144569-144712
35	329373	6682537	Minus	38950-39301
	329412	6682553	Minus	68948-69041
	329424	5868879	Plus	362196-362344
	329446	5868886	Plus	84776-84899
	329449	5868886	Plus	97697-97771

5

TABLE 14: shows genes, including expression sequence tags, down-regulated in prostate tumor tissue compared to normal prostate tissue as analyzed using Affymetrix/Eos Hu02 GeneChip array. Shown are the ratios of "average" normal prostate to "average" prostate cancer tissues.

10	Pkey: ExAccn: Unigene Unigene R1:	ID:	Exemplar A Unigene nu Unigene ge		
15	Pkey	ExAcen	UnigeneiD	Unigene Title	R1
15		AA281133 D60641	Hs.88808 Hs.131921		18.53 14.55
		Al251936	Hs.146298		12.17
00		AA418762	Hs.190044	ESTS	10.55
20		AA947682	Hs.217173	ESTs; Weakly similar to Chain A; Cdc42hs-Gdp Complex [H.saplens]	10.17
		AA543096 C05278	Hs.13648 Hs.166800	ESTs; Highly similar to mitogen-induced [M.musculus] ESTs; Moderately similar to [PYRUVATE DEHYDROGENASE(LIPOAMIDE)] KINASE ISOZYME 4 PRECURSOR [H.saplens]	9.2 8.87
	324882	AW419080	Hs.250645		8
25		U57796		zinc finger protein 192	7.88
	330126			CH.21_p2 gi 6093735	7.8
		AA737400	Hs.142230		7.7
		AA148950	Hs.188836	ESTs	7.64
	320668	R58399	Hs.146217	ESTs	7.4
30	330769	AA465192	Hs.16514	ESTs ·	7.15
	312614	AI766732	Hs.201194	ESTs	7
	314790	AW341754	Hs.189305	ESTs	6.83
	309979	AW452118	Hs.257533		6.74
~ -		AA743396	Hs.189023		6.49
35	329192			CH.X_hs gi 5868716	6.1
	-	AA627642	Hs.4994	transducer of ERBB2; 2 (TOB2)	5.99
		AW500106		EST duster (not in UniGene) with exon hit	5.82
		AW452382	Hs.257564		5.8
40		AA679001	Hs.192221		5.68 5.43
40		AA034364		ESTs; Weakly similar to !!!! ALU CLASS B WARNING ENTRY !!!! [H.sapiens]	5.43
		AW298141	Hs.157975		5.35
		F30712	11-440007	EST cluster (not in UniGene) with exon hit	5.31
		A1627358	Hs.148367		5.25
45		D84454	Hs.21899	CH22_FGENES.421_30	5.25
43	334719	AA813958	Ha 207727	ESTs; Moderately similar to KIAA0071 [H.sapiens]	5.22
		Al625304	Hs.190312		5.22
		AW511298	Hs.256067		5.19
		H86747		KIAA1116 protein	5.11
50	-	Al739168	1 ISLL I COL	EST cluster (not in UniGene)	4.97
-		AW206435	Hs.146057		4.97
	-	N50080	Hs.117827		4.78
		AW197887	Hs.253353		4.63
		AA766825		EST cluster (not in UniGene)	4.58
55	315850	AW270550	Hs.116957		4.53
	303124	AF161350		EST cluster (not in UniGene) with exon hit	4.46
	323346	AL134932	Hs.143607		4.4
	301383	AA913591	Hs.126480	ESTs	4.35
	324513	AW501678	Hs.164577		4.28
60	303480	AA331906		EST cluster (not in UniGene) with exon hit	4.25
		AA301270		EST cluster (not in UniGene)	4.22
		AW468119		EST cluster (not in UniGene)	4.2
		A1733395	Hs.129124	ESTs	4.1
		R42049	Hs.195473		4.08
65	-	AW451570	Hs.126850		4.03
	319750	AA621606	Hs.117956	ESIS	4.03

	000500	TEFORA		FOT chicken (act in HelCono)	4
		T55958	11- 404074	EST cluster (not in UniGene)	4
		AW026761	Hs.134374		4
		A1990652	Hs.208973		3.96
_		A1939339	Hs.146883		3.95
5		Al364186	11-057405	EST singleton (not in UniGene) with exon hit	3.94
		AW449204	Hs.257125		3.93
		W60843	Hs.31570		3.9
		Al679131	Hs.201424		3.88
10		AA541323	Hs.115831		3.83
10		N53442	Hs.143443		3.79
		AW203986	Hs.213003		3.75
		AA481027	HS.12/336	ESTs; Weakly similar to ORF YGR245c [S.cerevisiae]	3.74
		D59945	11. 420220	EST cluster (not in UniGene)	3.73
15		A1039702		collagen; type i; alpha 2	3.73 3.73
15		AA804409	Hs.118920		3.73
		AA765470	Hs.122826		3.68
		D81150	U- 454044	EST cluster (not in UniGene) with exon hit	3.66
•		Z38720	Hs.151014		3.59
20		AA011603	Hs.102332	EST cluster (not in UniGene)	3.52
20		H95082	ns. 102332		3.5
		AA631739 AA317561		EST duster (not in UniGene)	3.49
			Hs.115130	EST cluster (not in UniGene)	3.47
		AW516519	ns.113130	EST singleton (not in UniGene) with exon hit	3.46
25	300038	AW242630		AFFX control: MurlL4	3.38
43		Al088192	Un 125474	ESTs; Weakly similar to ATP-DEPENDENT RNA HELICASE A [H.sapiens]	3.36
		AA731520	Hs.170504		3.35
		AA196027		glyceraldehyde-3-phosphate dehydrogenase	3.34
		A1948688	Hs.191805		3.33
30	329815	A1340000	115.191000	CH.14_p2 gij6624888	3.32
50		A1745387	Hs.239124		3.31
		N53574	Hs.158932	· · · ·	3.3
	329218	1455574	113.100002	CH.X_hs gi 5868726	3.28
		AW440742	Hs.155556		3.28
35		AW503637	113.10000	EST cluster (not in UniGene) with exon hit	3.25
55		Al347274		EST singleton (not in UniGene) with exon hit	3.25
		AA324975	Hs.128993	ESTs; Weakly similar to KIAA0465 protein [H.sapiens]	3.25
		AA700439	Hs.188490	:	3.25
	334061			CH22_FGENES.327_14	3.23
40	336036			CH22_FGENES.678_7	3.23
		H67818	Hs.222059		3.21
		AW139383	Hs.245437	ESTs	3.2
	316733	AA811713	Hs.163222	ESTs	3.2
	300855	AW235248	Hs.79828	ESTs	3.2
45	323611	AA304986	Hs.145704	ESTs	3.19
	314138	AA740616		EST cluster (not in UniGene)	3.17
	316774	AA814859		EST cluster (not in UniGene)	3.16
	308884	AI833131	Hs.179100	ESTs	3.11
	331317	AA258222	Hs.87757	ESTs	3.1
50		A1989538	Hs.191074	ESTs	3.08
		AA749062	Hs.180285		3.08
	321040	H26953		EST duster (not in UniGene)	3.08
		A1824829		EST singleton (not in UniGene) with exon hit	3.08
ے ہے		AA236233	Hs.188716		3.07
55	316667	AW015940	Hs.232234		3.07
		AW503101		EST duster (not in UniGene)	3.07
		AW293046	Hs.255158		3.07
		Al239706	Hs.189886		3.06
60		AW297967	Hs.188181		3.05
60	312335	AW043620	Hs.236993		3.03 3.01
	322957	AA247755	Un 100700	EST duster (not in UniGene)	3.01
		AA830053	Hs.126798		2.99
		AA009660	ns.251948	ESTs; Moderately similar to T07D3.7 [C.elegans]	2.99
65		T27119	Un 104007	EST duster (not in UniGene)	2.97
UJ		A1076370	Hs.134037		2.96
		AA458637	Hs.152207		2.95
		AA053294 AW340014		EST singleton (not in UniGene) with exon hit EST singleton (not in UniGene) with exon hit	2.95
		H08035	He 21200	EST's ingleton (not in Unidene) with excert in. ESTs; Moderately similar to PUTATIVE GLUCOSAMINE-6-PHOSPHATE	2.00
	318001	110000	113-61330	Possi Minigarial & Similar in LAIVILLE appropriation LINE	

				ISOMERASE [H.sapiens]	2.95
	321253	A1699484		EST cluster (not in UniGene)	2.93
		AA149508	Hs.103288	· ·	2.93
	332864	751110000		CH22_FGENES.28_4	2.92
5	300027			VI UEI 120.20_1	
•	000021	M11507		AFFX control: transferrin receptor	2.91
	204220	AA884766		EST cluster (not in UniGene)	2.88
			Hs.170291		2.88
		AA137114	NS.170291		2.88
10	333916	710000		CH22_FGENES.296_5	2.87
10		Z43272		EST duster (not in UniGene)	2.87
		Al040125	Hs.150521		
	-	AA233056	Hs.191518		2.85
		AA825148	Hs.21229	F-box protein Fbw1b	2.84
	335862			CH22_FGENES.629_7	2.83
15	317672	AW205409	Hs.127748		2.82
	323416	AI610397	Hs.159560		2.81
	312652	Al419909	Hs.160994	ESTs	2.81
	324094	AA382603		EST cluster (not in UniGene)	2.81
	319761	R84237		EST cluster (not in UniGene)	2.8
20	317013	AA864468	Hs.135646	ESTs	2.8
	317383	AA913887	Hs.126511	ESTs	2.78
		AW277121	Hs.254881	ESTs	2.78
		Al950844	Hs.128738	ESTs; Weakly similar to non-lens beta gamma-crystallin like protein [H.sapiens]	2.77
	332808			CH22_FGENES.7_10	2.75
25		AW293826	Hs.250610		2.75
		C06003	Hs.116456	ESTs	2.73
		AW517542	Hs.208382		2.73
		AW296076	,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	EST singleton (not in UniGene) with exon hit	2.73
		AA683529	Hs.143119		2.73
30		AW294020	Hs.117721		2.72
50		AA062971	He 181161	ESTs; Weakly similar to INHIBITOR OF APOPTOSIS PROTEIN 1 [M.musculus]	2.72
		H53744	113.101101	EST cluster (not in UniGene)	2.72
		H77679		EST singleton (not in UniGene) with exon hit	2.72
		uttota		CH.13_hs gi 5866994	2.71
35	325602	DECOME	Hs.136698		2.71
22		R59096	ns.130090		2.71
		N75450	Un 150000	EST cluster (not in UniGene) with exon hit ESTs; Weakly similar to predicted using Genefinder [C.elegans]	2.69
		AA831215			2.68
		AI091458	Hs.134559		2.68
40		R38715		Homo sapiens clone 24540 mRNA sequence	2.68
40		AI823999	Hs.162000		2.67
		AA614308		EST singleton (not in UniGene) with exon hit	2.67
		AI431345	Hs.161784		
		AW193466	Hs.136525		2.67
40		AI057369	Hs.122536		2.65
45		AA135565	Hs.250739		2.65
		Al308989	Hs.156939		2.65
	324825	AA704457		ESTs; Moderately similar to gag [H.sapiens]	2.65
	300336	AW292417	Hs.255074	ESTs; Moderately similar to high-risk human papilloma viruses E6	
				oncoproteins targeted protein E6TP1 alpha [H.sapiens]	2.64
50	317850	N29974		EST cluster (not in UniGene)	2.64
	339047			CH22_DA59H18.GENSCAN28-7	2.64
	324580	AA492588		EST cluster (not in UniGene)	2.63
		Al817933	Hs.209584	ESTs	2.62
	319478	R06841		EST cluster (not in UniGene)	2.62
55 .	300793	Al248571	Hs.186837	ESTs	2.61
	313733	AA836116		EST cluster (not in UniGene)	2.6
	326505			CH.19_hs gi 5867435	2.6
		AW015506	Hs.130730		2.6
		AF090948		EST cluster (not in UniGene) with exon hit	2.59
60		H24244	Hs 240763	ESTs; Weakly similar to /prediction	2.58
		Al209108	Hs.143946		2.57
	329224	. 11200100	1101170070	CH.X_hs gi 5868728	2.56
	328018			CH.06_hs gi 5902482	2.56
		AA324437	Hs.177230		2.55
65		AW157377	Hs.132910		2.55
55		AW136134	Hs.220277		2.55
					2.54
		AI479011	Hs.170783		2.54
•		AJ743261 AW293174	Hs.131860 Hs.252627		2.53

	313966	Al807551	Hs.189061		2.53
	331263	AA015718		ze31a12.s1 Soares retina N2b4HR Homo sapiens cDNA clone	
				IMAGE:36574 3', mRNA sequence	2.51
~		AW055233	Hs.160870		2.5
5		AA085996	Hs.248572	Human PAC clone DJ404F18 from Xq23	2.5
		AJ001408		EST duster (not in UniGene) with exon hit	2.5
		Al613519		EST singleton (not in UniGene) with exon hit	2.49
		AF086538		EST cluster (not in UniGene)	2.49
10		AA974253	Hs.120319		2.49
10		AA203415	Hs.136200		2.48 2.48
		W76005	Hs.32094		2.48
	-	AA243617			2.47
		AA256675	H\$.200438	ESTs; Weakly similar to atypical PKC specific binding protein [R.norvegicus]	2.47
15	328744	Al624497		EST singleton (not in UniGene) with exon hit CH.07_hs gil5868290	2.47
15		W45574	Hs.252497		2.47
	328121	11400/4	113.232437	CH.06_hs gi[5868031	2.47
		A1670955	Hs.200151		2.46
		AA521381	Hs.187726	·	2.45
20		AA188868		ESTs; Weakly similar to NUCLEAR FACTOR 1/X [H.sapiens]	2.45
	329454			CH.Y_hs gi 5868887	2.45
	336605			CH22_FGENES.420_4	2.45
		AI444628	Hs.256809		2.44
	323362	AL135067	Hs.117182	ESTs	2.44
25	300024	M10098	AFFX contr	ot: 18S ribosomal RNA	2.44
	325026	AI671168	Hs.12285	ESTs	2.43
	324510	Al148353	Hs.120849		2.43
		A1765182	Hs.119903		2.43
~~		M78276	Hs.255917		2.43
30		AA041455	Hs.209312		2.43
		AW135854	Hs.132458		2.42
		R01342	11-005504	EST duster (not in UniGene)	2.42 2.42
		A1744361	HS.205591	ESTs; Weakly similar to zinc finger protein Png-1 [M.musculus]	2.42
35	327871			CH.06_hs gij5868131 CH22_FGENES.565-3	2.41
<i>JJ</i>	337173	AA465635		EST cluster (not in UniGene) with exon hit	2.41
		AL118754		EST cluster (not in UniGene)	2.4
		Al791138	Hs.116768		2.4
		AA830515	Hs.222917		2.4
40		AA253351		STAT induced STAT inhibitor-4	2.4
		AA017595		ESTs	2.4
		Al701559		EST singleton (not in UniGene) with exon hit	2.39
	313035	N36417	Hs.144928		2.37
	322114	AA643791	Hs.191740	ESTs	2.37
45		W49823	Hs.145553		2.37
	303211	AA099548	Hs.191436	ESTs; Highly similar to dJ1118D24.4 [H.sapiens]	2.37
		AA932948		EST cluster (not in UniGene) with exon hit	2.36
	338165			CH22_EM:AC005500.GENSCAN.212-3	2.36
50		AA557952	11	EST duster (not in UniGene)	2.35 2.35
50	-	AA779704	Hs.168830		2.35
		R41582		retinal degeneration B beta	
		T48446	Hs.193162		2.35 2.35
		AA663726 AA286678	Hs.116922	EST cluster (not in UniGene) with exon hit	2.34
55		AW292740	Hs.254815		2.34
55		AA682305	Hs.133268		2.33
		AA642912	113.100200	EST singleton (not in UniGene) with exon hit	2.33
		Al000320		EST singleton (not in UniGene) with exon hit	2.33
		Al651016	Hs.246311		2.33
60	321348	Z49979		EST cluster (not in UniGene)	2.33
	325112	Al903770	Hs.124344		2.32
	336679			CH22_FGENES.43-7	2.32
	321383	AJ002574		EST cluster (not in UniGene)	2.32
	337357			CH22_FGENES.730-6	2.31
65	300680	AW468066	Hs.257712	ESTs; Weakly similar to KIAA0986 protein [H.sapiens]	2.31
	327120			CH.21_hs gi[6531970	2.31
		AW250553		EST duster (not in UniGene) with exon hit	2.3
		Al475490	Hs.170577		2.3 2.3
	215520	AAR27652		FST duster (not in UniGene)	2.3

	312189	T95594	Hs.187435	ESTs	2.3
		AA991705		EST singleton (not in UniGene) with exon hit	2.3
	327061			CH.21_hs gij6531965	2.3
		AA759098	Hs.192007		2.3
5		Al968646	Hs.33862		2.29
9		AA203339	Hs.220750		2.29
		A1680915	Hs.201379		2.28
	335250	A1000313	115.201373	CH22_FGENES.516_11	2.28
		700007	Un 01660		2.28
10		Z38907		KIAA0888 protein	2.28
10		AW294013	Hs.200942	·	2.28
		AA969121	Hs.254296		2.28
		A1608881		ESTs; Highly similar to junctional adhesion molecule [H.saplens]	2.28
		Al970543	Hs.192605		2.28
15		Z43395	11: 404000	EST duster (not in UniGene)	2.27
15		AA252753	Hs.164039		
		AA342250		ubiquitin specific protease 16	2.27 2.27
		AW292127	Hs.144758	·	
		AA766025	Hs.238794		2.27
20		Al697668	Hs.202241		2.26
20 .		AA229781	Hs.221962		2.26
		Al004614	Hs.130577		2.26
		AW474196	Hs.221604		2.25
		AA769123		EST cluster (not in UniGene)	2.25
٥.		AA968799	Hs.150289	· = : · · · · · · · · · · · · ·	2.25
25		AA330095		EST cluster (not in UniGene)	2.25
		A1000929		EST singleton (not in UniGene) with exon hit	2.24
	329109			CH.X_hs gi 5868626	2.24
	311043	A1871209	Hs.177128		2.24
••		Al458372		ESTs; Weakly similar to synapsin lb [M.musculus]	2.24
30		Al193698	Hs.184776	ribosomal protein L23a	2.24
		A1888045		EST singleton (not in UniGene) with exon hit	2.23
	310749	A1493675	Hs.170332		2.23
		Al914939	Hs.212184		2.22
	320409	AA356195		EST cluster (not in UniGene)	2.21
35	333149			CH22_FGENES.87_8	2.21
	324951	M86125	Hs.137487		2.21
	321939	A1791617	Hs.145068	ESTs	2.2
	320594	Al863952		arginyftransferase 1	2.2
	320722	R67430	Hs.172787		2.2
40	321781	D78667		EST cluster (not in UniGene)	2.2
	328903			CH.08_hs gi 5868514	2.2
	303889	T19204		EST duster (not in UniGene) with exon hit	2.2
	325045	T08845		EST duster (not in UniGene)	2.2
	312828	A1865455	Hs.211818	ESTs; Moderately similar to IIII ALU SUBFAMILY J WARNING ENTRY IIII [H.sapie	
45	335109			CH22_FGENES.494_15	2.18
		AA131471	Hs.71440		2.18
		Al971362	Hs.231945		2.18
		AA513456		EST singleton (not in UniGene) with exon hit	2.18
70	337393			CH22_FGENES.747-4	2.18
50	332812			CH22_FGENES.7_14	2.18
	327665			CH.04_hs gij5867839	2.18
		AW504859	Hs.237849		2.17
	326508			CH.19_hs gi 6682496	2.17
		AW161535	Hs.258803		2.17
55		Al765651	Hs.172900		2.17
		AW276810	Hs.254859		2.16
	311179	AI880843	Hs.223333		2.16
	315320	AI084182	Hs.186895		2.16
		AI015203	Hs.118015		2.16
60		AW139117	Hs.117494		2.15
	300864	AA406539	Hs.190958		2.15
		AA463262		EST cluster (not in UniGene)	2.15
	322574	AF156548		EST cluster (not in UniGene)	2.15
		C03864		EST cluster (not in UniGene)	2.15
65		AA002047		EST cluster (not in UniGene).	2.14
		AA353895	Hs.152983	HUS1 (S. pombe) checkpoint homolog	2.14
	337646		•	CH22_EM:AC000097.GENSCAN.11-2	2.13
		AF174008		EST cluster (not in UniGene) with exon hit	2.13
	312185	AA654772	Hs.186564	ESTs	2.13

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					0.40
		A1066544 AA602917	Un 150074	EST singleton (not in UniGene) with exon hit	2.13 2.12
		AA602917 Al821782	Hs.156974	ESTS; Moderately similar to IIII ALU SUBFAMILY SC WARNING ENTRY IIII [H.sap	
		Al800041	Hs.190555		2.11
5		R66867		EST cluster (not in UniGene)	2.11
		Al167877	Hs.143716		2.11
		AW015206	Hs.178784		2.11
		AA235482		ferritin; heavy polypeptide 1	2.11 2.1
10		AA399018 T72744	Hs.250835	EST cluster (not in UniGene)	2.1
10	328078	112144		CH.06_hs gij5868008	2.1
		AW090770	Hs.192271		2.1
		Al738720		EST singleton (not in UniGene) with exon hit	2.09
15		AW439969	Hs.218177		2.09
15		AI761786	Hs.204674		2.09 2.08
	332933	AA848118	Hs.221216	CH22_FGENES.38_7	2.08
	325498	•		CH.12_hs gij5866967	2.08
		AW296067	Hs.124106		2.08
20		AW149321	Hs.105411		2.08
		AA640770		EST cluster (not in UniGene)	2.07
		AA347452		EST duster (not in UniGene) with exon hit	2.07
		AW450674	Hs.114696		2.06 2.06
25	326920 327574			CH.21_hs gl 6456782 CH.03_hs gl 5867818	2.06
25		Al052795	Hs.192201		2.06
		AW503733	Hs.170315		2.05
	305235	AA670480		EST singleton (not in UniGene) with exon hit	2.05
		AA693880		EST cluster (not in UniGene)	2.05
30		AW445167	Hs.126036		2.05
		AW408683	Hs.32922		2.05 2.05
	335146	Al678183	He 170017	CH22_FGENES.499_2 prostaglandin E receptor 3 (subtype EP3)	2.04
		AA120970	Hs.143199		2.04
35		R62925	Hs.243665		2.04
		AA290875	Hs.30120		2.04
		Al215643	Hs.171381		2.03
		W23285		EST duster (not in UniGene)	2.03
40		AA282197	Hs.89002	ESTs; Highly similar to CGI-07 protein [H.sapiens]	2.03 2.03
40		AA994530 Al298794	Hs.129130	EST singleton (not in UniGene) with exon hit	2.03
		Al493742	Hs.165210		2.02
		AW294522	Hs.149991		2.02
		AW245528	Hs.134754		2.02
45	331286	AA137062	Hs.103853		2.01
		Al989942	Hs.232150		2.01
	335601	Alconono	11-004074	CH22_FGENES.581_41	2.01 2.01
,		Al682303 AA249018	Hs.201274	EST cluster (not in UniGene)	2.01
50	328190	AA243010		CH.06_hs gij5868077	2
-	338030			CH22_EM:AC005500.GENSCAN.148-16	2
	333940			CH22_FGENES.301_6	2
	328227	•		CH.06_hs gi 5868105	2
55		N27448	Hs.43944	EST	2
55	335288	A1074007		CH22_FGENES.527_1	2 2
		Al274307 AL134620		EST singleton (not in UniGene) with exon hit EST cluster (not in UniGene)	2
		R21945	Hs.256153	·	2
		AA502583	Hs.197271		2
60	327489			CH.02_hs gi 6004459	1.99
		AW175841	Hs.192183		1.99
		AW168096	Hs.195188	glyceraldehyde-3-phosphate dehydrogenase	1.99
	337043	A1000474	U- 007040	CH22_FGENES.439-19	1.98 1.98
65		Al828174 Al370434	Hs.227049	EST singleton (not in UniGene) with exon hit	1.98
05	328656	71010404		CH.07_hs gi 6004473	1.98
		AA813784	Hs.123001		1.98
		W45302		helicase-mol	1.98
		AA701499	Hs.148115		1.98

	313171	N67879	Hs.157695	ESTs	1.97
		Al241421	Hs.132236	ESTs	1.97
		N66393	Hs.102754		1.97
			Hs.226803		1.97
5		Al962180			1.97
J	335864			0,122,,120,000_0	1.97
		W00545	Hs.171785	2010	
	314065	AA868267	Hs.85524	2010	1.96
	323086	H15474	Hs.12214	1 tottle capitatic cione per 10 mi national conse	1.96
	323919	AA862973	Hs.220704	ESTs	1.96
10		Al373163	Hs.170333		1.96
10		AW090537	110.11.0000		1.96
				Edit director fraction derivative, with annual con-	1.96
		AW028820	11- 000004	20; distribution of the contract of	1.95
		Al820675	Hs.203804		
	323787	AW373446	Hs.169885	ESTs; Weakly similar to cDNA EST EMBL:T02216 comes from this gene [C.elegans	
15	338112			O. M	1.95
	313625	AW468402	Hs.254020	2010	1.95
•	325240			CH.10_hs gi 5866848	1.95
	331833	AA412102	Hs.250911	interleukin 13 receptor, alpha 1	1.95
		N63882		za21f9.s1 Soares tetal liver spleen 1NFLS Homo sapiens cDNA clone	
20	OULLUL	1100002			1.95
20	000070	A14/007/07	U- 052017	mb received of man and and	1.95
		AW237425	Hs.253817	20.0	1.95
	326023				
	321609	H86021		ESTs; Weakly similar to hMmTRA1b [H.sapiens]	1.94
	324183	AA402453	Hs.113011	.ESTs	1.94
25	336276			CH22_FGENES.762_5	1.94
	334913			CH22 FGENES.456_3	1.94
	325417			CH.12_hs gl 5866925	1.94
		AW043590	Hs.225023		1.94
			113.223020		1.94
20		Al148763		DOT DISCOTT (COTTO CINCOTT)	1.94
30		Al092235		EST singleton (not in UniGene) with exon hit	1.94
		AW452948	Hs.257631		
	321289	R84687	Hs.226306		1.94
	308521	A1689808		EST singleton (not in UniGene) with exon hit	1.93
	306382	AA968967		EST singleton (not in UniGene) with exon hit	1.93
35		AA262999	Hs.42788		1.93
-		AA501412		ESTs; Weakly similar to Pro-Pol-dUTPase polyprotein [M.musculus]	1.93
		AW168753	110.101000	EST singleton (not in UniGene) with exon hit	1.93
		A11 100/33		CH.21_hs gi 5867664	1.93
	327014	414/005000		EST cluster (not in UniGene) with exon hit	1.93
40		AW025860			1.92
40		AA995223	Hs.129559	ESI	1.92
		AA019806		spinocerebellar ataxia 7 (olivopontocerebellar atrophy with retinal degeneration)	
	313083	N50545	Hs.159200	ESTs	1.92
	327752			CH.05_hs gij5867949	1.92
	318674	AA295490		EST cluster (not in UniGene)	1.92
45		AW297762	Hs.255690		1.91
		AA608787	Hs.112590		1.91
		AL036947	113.112330	EST duster (not in UniGene)	1.91
					1.91
		AA317554	11. 000.00	EST duster (not in UniGene)	1.91
50		AI765013	Hs.209128		
50	300976	Al246374	Hs.185861		1.91
	323715	AA322155		EST duster (not in UniGene)	1.91
	313800	AW296132	Hs.166674	ESTs	1.91
		AA489697	Hs.145053	ESTs .	1.91
		AW518573		Immunoglobulin kappa variable 1D-8	1.91
55		AA354549	He 41181	Homo sapiens mRNA; cDNA DKFZp727C191 (from clone DKFZp727C191)	1.91
<i></i>			110.41101	CH22_FGENES.339_1	1.9
	334150		Un 225040		1.9
		AW450967	Hs.235240		1.9
		AW207642	Hs.174021		
	324774	AI031771	Hs.132586	77.7	1.9
60	326507			CH.19_hs gi 5867435	1.9
		AA405696		EST cluster (not in UniGene)	1.9
	336268			CH22_FGENES.758_2	1.9
	915970	Al985544	Hs.116429	-	1.9
					1.9
45	325824		Un 040000	CH.15_hs gi 5867048	1.9
65 ·		AA737780	Hs.213392		1.9
		AA418583	Hs.143621		
		AA961643	Hs.127716	*	1.89
	307050	Al147341	Hs.146734	EST	1.89
	306830	AI075803		EST singleton (not in UniGene) with exon hit	1.89

		AL049925			1.89
	320127	H72615	Hs.17268		1.89
	337736				1.89
_	331319	AA262755	Hs.194264	ESTs	1.88
5	310767	Al377505	Hs.158835		1.88
	314880	A1732169	Hs.105429	ESTs	1.88
	312539	Al004377	Hs.200360		1.88
	309674	AW205604	Hs.168034	ESTs; Weakly similar to !!!! ALU SUBFAMILY SP WARNING ENTRY !!!! [H.sapiens]	1.88
	314621	Al627478	Hs.187670		1.88
10		Al972146	Hs.192756		1.88
		AA007374			1.88
		U09060			1.88
	329511	000000		CH.10_p2 gi[3983514	1.88
		AI699412	Hs.201925	armina Par Bilanasa	1.87
15		AI815985			1.87
10		AA725670	He 120485	ESTs; Weakly similar to serine/threonine kinase with SH3 domain; leucine	
	001100	MILLOID	113.120100		1.87
	332222	AI20271	Hs.176618		1.87
	-	AA055475			1.87
20		Al159863	Hs.143713		1.87
20		AW291847		2010	1.87
			113.121713	manual commence on bionemili marketing	1.86
		Al827817	U- 10000		1.86
		R84768	ms. 10088	Treme deplete didite address in the conference	1.86
25	325587	41004040	11-450000	ar manager and a financial control of the control o	1.86
23		Al884313	Hs.158906		1.86
		R13085		Tot Cloth (noth on the one)	1.86
		AA317915			1.86
	338427			01122_2M3 1000000102.11010 1	
20		Al352293	Hs.191098		1.85
30 .		H85330	Hs.146060		1.85
		F05865	Hs.249180	and any confederal control and any control and any	1.85
		AJ230822		To condition (not in otherway) that should	1.85
		Al679966	Hs.150603	2010	1.85
25		Al239811	Hs.157491		1.85
35		AW016437	Hs.233462		1.84
		AA278347	Hs.126078		1.84
	335586				1.84
	339209			01.22C11.1021110211001110	1.84
	307954	Al419692		Tot on Broth from the order	1.84
40	302549	AF055136		To de constitue de la constitu	1.84
		H87213	Hs.158092	20.0	1.84
	301239	AA807558		Tot care (not in one one) that of	1.84
	332434	N75542	Hs.75356	adiosipaeti taetei i	1.84
	327192	•		4: 110 1_110 Bilecot : 10	1.83
45	310214	A1220072	Hs.165893	ESTs	1.83
	320516	R33857	Hs.181479	ESTs; Weakly similar to E-SELECTIN PRECURSOR [H.sapiens]	1.83
	324231	W60827		EST cluster (not in UniGene)	1.83
	336616			CH22_FGENES.613_5	1.83
	328799			CH.07_hs gi 5868316	1.83
50	324661	AW504161		EST cluster (not in UniGene)	1.83
	313190	AA766707	Hs.153039		1.83
	301979	L28168	Hs.121495	potassium voltage-gated channel; Isk-related family; member 1	1.82
	302099	AL021397	Hs.137576	ribosomal protein L34 pseudogene 1	1.82
	320187	T99949		EST cluster (not in UniGene)	1.82
55	320791	R78808	Hs.93961	ESTs; Weakly similar to !!!! ALU CLASS A WARNING ENTRY !!!! [H.sapiens]	1.82
	305733	AA829535	Hs.84298	CD74 antigen (invariant polypept of MHC; class II antigen-associated)	1.82
		AI569349		ribosomal protein S9	1.81
		W78877	Hs.40111		1.81
		Al915122		ESTs; Weakly similar to F33D11.9b [C.etegans]	1.81
60		H90265	Hs.100636		1.81
	329519	•••	*	CH.10_p2 gi 3983510	1.81
		AA220982		EST cluster (not in UniGene)	1.81
		N62937	Hs.139181		1.81
	329246			CH.X_hs gij5868732	1.81
65		AA481271	Hs.193945	,	1.81
-		A1420990	Hs.161303		1.81
	325866			CH.16_hs gi 5867076	1.81
		Z78343		EST cluster (not in UniGene)	1.8
	333712			CH22_FGENES.251_1	1.8

	313457	AA576052	Hs.193223		8.1
	321591	H85687	Hs.117927	ESTs 1	1.8
	330260			CH.05_p2 gi 6671884	8.1
	311080	A1656320	Hs.197711	ESTs 1	8.1
5	329522			CH.10_p2 gi[3983507	8.1
-		AA081924	Hs.211417	~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~	8.1
		Al275011	Hs.204877		8.1
		H20560	Hs.244624	and the second s	1.8
		Al341180		40 .0	1.79
10			113.130113	and the state of t	1.79
10		R17531	11- 400004		1.79
		AA730673	Hs.188634		1.79
		AI400310	Hs.148958	##··	1.79
		AW292760		The state of the s	
1 =	326506				1.79
15 [,]		AA649011	Hs.187902		1.79
		AI623739	Hs.186387	20.0	1.79
	312180	Al248285	Hs.118348		1.79
	313058	D81015	Hs.125382		1.79
	330120			0.1p_ 6/jour. 100.	1.78
20	328412				1.78
	302345	NM_000565		mot states (not at states)	1.78
	308100	Al475949		EST singleton (not in UniGene) with exon hit	1.78
	311386	AW205705	Hs.207514	ESTs	1.78
	330282			CH.05_p2 gi[6671910	1.78
25		Z43011	Hs.21169		1.78
25		AA845630	Hs.117904		1.78
	325450				1.78
		H54178	Hs.226469		1.78
		H20826		E010	1.78
30		AA333666	113.51705	2010	1.77
20		Al264671	Hs.164166	and the state of t	1.77
					1.77
		AI540166	Hs.129563 Hs.128245		1.77
		Al683782	Ho 100240	ESTs; Weakly similar to !!!! ALU SUBFAMILY SB WARNING ENTRY !!!! [H.sapiens]	
35		AL038841	HS. 100010		1.77
33	336123	41000400	U= 000404	o	1.77
		Al286182	Hs.208484		1.77
		AW451733	Hs.119824		1.77
		AA001811	Hs.83722		1.77
40	329941			or a rough gap root of	1.77
40	328329				
		Al493054	Hs.158968	40.0	1.77
	325902			3.1.1.2.1.0 8,1000.101	1.76
		W01813		The topical time.	1.76
		A1274851	Hs.258744	20.0	1.76
45		A1025527	Hs.222097		1.76
	331909	AA437300	Hs.178210	44.0	1.76
	321553	H92449	Hs:116406	2010	1.76
	301618	T52760		and the state of t	1.76
	319592	AA627356	Hs.163315		1.76
50	318511	T26528	Hs.227175	ESTs; Weakly similar to !!!! ALU SUBFAMILY SQ WARNING ENTRY !!!! [H.sapiens]	1.76
	327183				1.76
	313516	AA029058	Hs.135145		1.76
		- Al752482		EST cluster (not in UniGene)	1.76
		AA419617		EST cluster (not in UniGene)	1.76
55		AW451142	Hs.255628		1.76
		AW449374	Hs.257149	ESTs	1.75
	319775	AA504429	Hs.6211	methyl-CpG binding domain protein 1	1.75
	314775	AJ149880	Hs.188809		1.75
	337460	74110000	113.100000		1.75
60		AW297444			1.75
UU		AA995014	Uc 120544	ESTs; Weakly similar to ORF YLL027w [S.cerevisiae]	1.75
	040700	AM333014			1.75
	312/39	Al318426	Hs.155925	40.0	1.75
		H15355	Hs.60887		1.75
65	326495				1.75 1.75
65	337497		11- 4-6		1.75 1.75
	322633	AA004534	Hs.153981		
		F10812	Hs.101433	20.0	1.75
	326930				1.75
	216002	A AR37332		EST dueter (not in I IniGena)	1.75

		•			
	324826	AA704806	Hs.143842	ESTs	1.75
	311269	Al656924	Hs.174257	ESTs	1.75
	309375	AW075342		EST singleton (not in UniGene) with exon hit	1.75
	314171	AI821895	Hs.193481		1.75
5		Al990741	Hs.252809	ESTs	1.75
- 3	334387			CH22_FGENES.380_1	1.75
	-	Al300101	Hs.252222		1.75
		Al418055	Hs.161160		1.74
		AW501470	113.101100	EST duster (not in UniGene)	1.74
10		Al762929	No 206124	ESTs; Weakly similar to similar to reverse transcriptase [C.elegans]	1.74
10			115.200154	EST singleton (not in UniGene) with exon hit	1.74
		AW339340		EST duster (not in UniGene) with exon hit	1.74
		AW501336		EST cluster (not in UniGene)	1.74
		Z24981			1.74
15		AF111178		EST duster (not in UniGene) with exon hit	
15		W92924	11-0000	EST cluster (not in UniGene)	1.74
		H93199	Hs.33665		1.74
		AA737345		EST duster (not in UniGene)	1.74
	332243	N55484	HS.220540	ESTS; Highly similar to ARYL HYDROCARBON RECEPTOR NUCLEAR	4 74
20				TRANSLOCATOR [H.sapiens]	1.74
20		H02566		Homo sapiens mRNA; cDNA DKFZp434N174 (from clone DKFZp434N174)	1.74
		AL045752	Hs.211519		1.73
	320630	AA199847		EST cluster (not in UniGene)	1.73
	327288			CH.01_hs gi 5867481	1.73
	314986	Al201367	Hs.142860	ESTs	1.73
25	319078	H17255	Hs.144515	ESTs	1.73
	326278			CH.17_hs gij5867269	1.73
	302552	H49792		EST cluster (not in UniGene) with exon hit	1.73
	322322	AF086431		EST duster (not in UniGene)	1.73
	327075			CH.21_hs gi 6531965	1.73
30		Al797588	Hs.145459	ESTs	1.73
	300810	AI076890	Hs.186949	ESTs	1.73
		AA830893	Hs.119769		1.73
		AA773580	Hs.193598	ESTs	1.73
		AA004699		putative translation initiation factor	1.73
35		AW296802	Hs.255580		1.73
-		AI689617	Hs.200934		1.73
		F09774	Hs.175971		1.73
		Al984592	Hs.15088		1.73
		AA663560		ESTs; Weakly similar to K11C4.2 [C.elegans]	1.73
40		AW303457	11320075	EST duster (not in UniGene)	1.72
TU .		T71739	Hs.75442		1.72
		AI033922	Hs.122517		1.72
	334379	A1033322	113.122317	CH22_FGENES.379_11	1.72
		AA862733		EST singleton (not in UniGene) with exon hit	1.72
45			Hs.186566		1.72
40	329728	N34927	ns.100000		1.72
		METERO	Un 110064	CH.14_p2 gi 6065785	1.72
		N57692	Hs.118064 Hs.192386		1.72
		AL134875			1.12
50	302077	AA310580	ns. 132090	Homo sapiens chromosome 11; BAC CIT-HSP-311e8 (BC269730)	1.71
20	040700	A1074 400	11- 450004	containing the hFEN1 gene	1.71
		Al971438	Hs.158824		
		Al809985	Hs.203340		1.71
		AW238064	Hs.253909		1.71
55		H71999	11-044500	EST duster (not in UniGene)	1.71
55		T78791	Hs.241569	ESTs; Moderately smir to IIII ALU SUBFAMILY SQ WARNING ENTRY !!!! [H.saple	
		R56151		EST cluster (not in UniGene) with exon hit	1.71
	329089			CH.X_hs g 5868614	1.71
		AF086467		EST cluster (not in UniGene)	1.71
-60		Al080361	Hs.134217		1.71
60		AA489792		EST singleton (not in UniGene) with exon hit	1.71
		Al028149		pyruvate dehydrogenase kinase; isoenzyme 3	1.71
	310250	Al478629	Hs.158465		1.71
	338178			CH22_EM:AC005500.GENSCAN.219-6	1.71
	338910			CH22_DJ32I10.GENSCAN.11-2	1.71
65	321225	AL080073	Hs.251414	Homo sapiens mRNA; cDNA DKFZp564B1462 (from clone DKFZp564B1462)	1.7
	322289	AA534550	Hs.539	ribosomal protein S29	1.7
	319802	AI701489	Hs.202501	ESTs	1.7
	314022	AW452420	Hs.248678	ESTs	1.7
		AA515602	Hs.152330		1.7

	000000	8.8704000	11-000000	ECT-		1.7
		AA761322	Hs.220538			1.7
	304398	AA262785		EST singleton (not in UniGene) with exon hit		
	313421	AW339515	Hs.163700	ESTs		1.7
	309763	AW270182		EST singleton (not in UniGene) with exon hit	•	1.7
. 5		AF085833		EST cluster (not in UniGene)		1.7
,			Hs.121158			1.7
		AA764768				1.7
		T08597		EST cluster (not in UniGene)		
	327157			CH.01_hs gi 5866841		1.7
	. 314809	A1741461	Hs.161904	ESTs	· ·	1.7
10		H67220	Hs.146406		•	1.69
10			Hs.43616			1.69
		AW402302	118.43010			1.69
	328624			CH.07_hs gi[5868246		
	303344	AA255977	Hs.250646	ESTs; Highly similar to ubiquitin-conjugating enzyme [M.musculus]		1.69
	328960			CH.08_hs gi[6456775		1.69
15	315702	AA657501	Hs.146315	ESTs		1.69
		AJ224172	He 204096	lipophilin B (uteroglobin family member); prostatein-like		1.68
		R14537	110201000	EST cluster (not in UniGene)		1.68
				EST singleton (not in UniGene) with exon hit		1.68
		AW137700				1.68
	330417	D84424		hyaluronan synthase 1		
20	315296	AA876905	Hs.125286			1.68
	328538			CH.07_hs gi 5868485		1.68
		AA354146		EST cluster (not in UniGene)	_	1.68
		AL079289	He 137154	Homo sapiens mRNA full length insert cDNA clone EUROIMAGE 35971	_	1.68
			Un 1100E2	ESTs; Weakly similar to R10D12.12 [C.elegans]		1.68
25		Al927068				1.68
25		Al472124	Hs.157757			
	307512	Al273815	Hs.242463	keratin 8		1.68
	338506			CH22_EM:AC005500.GENSCAN.390-10		1.68
	331722	AA195405	Hs.110347	Homo saplens mRNA for alpha integrin binding protein 80; partial		1.68
		R05385		EST cluster (not in UniGene) with exon hit		1.68
30		Z42977	Hs.21062			1.68
J 0						1.68
		AW244073	Hs.145946			1.68
		AW137772	Hs.185980			
	325780			CH.14_hs gi 6381953		1.67
	321739	AL080280		EST duster (not in UniGene)		1.67
35	319808	T58960		EST cluster (not in UniGene)		1.67
	313443	AA249037		EST cluster (not in UniGene)		1.67
		AA424754	Hs.43149			1.67
		AI797592	Hs.207407			1.67
			115.207407			1.67
40 .	-	AA081820		EST duster (not in UniGene)		1.67
40	330320			CH.08_p2 gi[5932415		
	329081			CH.X_hs gi[5868602		1.67
	334026			CH22_FGENES.318_3		1.67
	317791	AJ801500	Hs.128457	ESTs		1.67
		AF086106		EST duster (not in UniGene)		1.66
45		R73816	Hs.17385			1.66
43		UISOID	HS. 17 303			1.66
	325452			CH.12_hs gi 5866941		1.66
	315106	AW452184	Hs.232100	and the second s		
	326014			CH.16_hs gi 5867160		1.66
	307130	Al185234		EST singleton (not in UniGene) with exon hit		1.66
50	300943	AA524545	Hs.224630	ESTs		1.66
		W21298		EST cluster (not in UniGene)		1.66
			He 170/37	ESTs; Weakly similar to hyperpolarization-activated; cyclic		
	310003	Al457946	115.170457	nucleotide-gated channel 2 [H.sapiens]		1.66
	323371	AL135118		EST cluster (not in UniGene)		1.66
55	335568			CH22_FGENES.581_4		1.66
	320654	AW263086	Hs.118112	ESTs		1.66
	338983			CH22_DA59H18.GENSCAN.3-1		1.65
	330002			CH.16_p2 gij6623963		1.65
			Hs.179891			1.65
60		AW205477	H2.1/2021			1.65
60	334487			CH22_FGENES.395_9		
		Al064824	Hs.193385			1.65
	309668	AW204480	Hs.253414	EST		1.65
	309518	AW148928	Hs.248895	EST		1.65
		Al421641		EST singleton (not in UniGene) with exon hit		1.65
65		AW369770	Hs.130351			1.65
J						1.65
		AA401858	Hs.224843			1.65
	338763			CH22_EM:AC005500.GENSCAN.517-16		
		AA232729	Hs.154302			1.65
	313231	AW139993	Hs.163682	ESTs		1.65

	334073			O. 1127. C. 11. 12. 12. 12. 12. 12. 12. 12. 12. 12	1.65
	319901	T77136	Hs.8765	Tita (nonecos romas promin	1.65
	326530			CH.19_hs gi 5867441	1.65
	301126	AI802877	Hs.210843	ESTs; Weakly similar to dJ1039K5.2 [H.saplens]	1.65
5		AA827082			1.65
-		AA236027			1.65
		AA099732			1.65
		AA033132		TO I MODIO! (MODING)	1.64
	337272	4.4000700	11-040004	O' ELL_1 GENERAL T	1.64
10		AA262768	HS.243901	turation protein	1.64
10		Z44266	11.050440	20. orașie, fried in amazire,	1.64
		AW342028	Hs.256112	20.0	
		AW293704	Hs.122658	2010	1.64
		AW295409	Hs.137945		1.64
	313835	AI538438	Hs.159087		1.64
15	320035	AA378974	Hs.130720	ESTs; Weakly similar to CELLULAR NUCLEIC ACID BINDING PROTEIN [H.sapien:	
	309372	AW074330			1.63
	324157	AW402236		201 district friends of the control of	1.63
	323929	AA354940 .	Hs.145958	2010	1.63
	302490	AA885502	Hs.187032	ESTs	1.63
20	333942			CH22_FGENES.301_8	1.63
	327469			CH.02_hs gi 5867772	1.63
		AA476777			1.63
		AI744068	Hs.160712		1.63
		AA282572			1.63
25		Al341594			1.63
23		F11623	113.107000	COLO Lucation and an are because it mande and	1.63
		Al962234	Hs.196102	201 district fraction of the control	1.63
			Hs.831	3-hydroxymethyl-3-methylglutaryl-Coenzyme A lyase (hydroxymethylglutaricaciduria	
		A1348076	П5.031	EST singleton (not in UniGene) with exon hit	1.63
20		AA989230	11- 400000	Edit Oligioton (not in discours) mai onem mi	1.63
30		AA086110	TIS. 188530		1.63
		Al269069		CO10, 1 18 11 Oct men to endergrand and analysis of market and	1.63
	328291		,	Ot 1:01_1:0 8-facecood	
		W93278		Co. Cargotta (not an one care)	1.63
~~		Al791700	Hs.127893	2010	1.63
35		AW440133	Hs.189690		1.62
	312834	AI028309	Hs.114246	LOID	1.62
	325326			C11.11_10 8. second	1.62
	313663	Al953261	Hs.169813	2010	1.62
	327526				1.62
40	300429	AW449679	Hs.156739	Colol tuling outlier to see on the time to the color of the color of	1.62
	305169	AA663131		EST singleton (not in UniGene) with exon hit	1.62
	316621	Al021996	Hs.122138		1.62
	329666			CH.14_p2 gij6272129	1.62
		Al744130	Hs.131201		1.62
45		AL031709			1.62
		Al307229	Hs.184304		1.62
		AA496019	Hs.201591	2010	1.62
		Al183686	113201331		1.62
					1.62
50		N49476	Hs.33439	ESTs; Weakly similar to IIII ALU SUBFAMILY J WARNING ENTRY IIII [H.sapiens]	
20		R87650	U2'''	CH22_EM:AC005500.GENSCAN.246-9	1.61
	338220	*****	11- 440040	CM2Z_EMIACOODOO.GENDOAN.240-5	1.61
		AW515270	Hs.118342		1.61
		AA984133		0 021 minutes 3 kinsters	
سے سے		R28628	Hs.203669		1.61
55	304569	AA490934			1.61
	313179	Al076101	Hs.131704		1.61
	326858			At 1970 Prince 107	1.61
	317276	AI823847	Hs.129986	ESTs	1.61
	312572	AA350125	Hs.187499	ESTs	1.61
60		AW451654	Hs.257482	ESTs	1.61
		AA452310	Hs.26090	ESTs; Wealdy similar to T20B12.1 [C.elegans]	1.61
		Al636253	Hs.196511		1.61
		A1620617	Hs.148565		1.61
	337780			CH22 EM:AC000097.GENSCAN.121-2	1.61
65				CH.05_hs gij5867982	1.61
UJ.	327796			EST singleton (not in UniGene) with exon hit	1.61
		Al610791	He 19Econ		1.61
		AI378032	Hs.125892		1.61
		AA437414		EST duster (not in UniGene) with exon hit	1.61
	337884			CH22 EM:AC005500.GENSCAN.54-2	

	000000	AACOTEAC	U- 4404E4	FOT-	4.04
		AA397546	Hs.119151		1.61
	303481	AA336839		EST cluster (not in UniGene) with exon hit	1.61
	314481	AA548589	Hs.105846		1.61
		Al908894	Hs.245893		1.6
5			115240050		
J		AA262442		EST duster (not in UniGene)	1.6
	326154			CH.17_hs gl 5867170	1.6
	331920	AA446885	Hs.99087 -	ESTs; Moderately similar to ZINC FINGER PROTEIN 141 [H.sapiens]	1.6
		AW406878		EST duster (not in UniGene)	1.6
10		W56710		EST duster (not in UniGene)	1.6
10	310597	Al739071	Hs.158515	ESTs	1.6
	307871	Al368665		EST singleton (not in UniGene) with exon hit	1.6
		AF088005		EST duster (not in UniGene)	1.6
		Al139857	Hs.143837		
					1.6
4 ==	332217	H98987	Hs.102383	EST	1.6
15	324937	M79230	Hs.192398	ESTs .	1.6
	320543	AF052176	Hs.158529	Homo sapiens clone 24457 mRNA sequence	1.6
		AW467388		EST duster (not in UniGene) with exon hit	1.6
			11- 404700		
		Al241331	Hs.131765		1.6
	319713	R24204		EST duster (not in UniGene)	1.6
20	301210	A1379982	Hs.158944	ESTs	1.6
		AW072861		EST singleton (not in UniGene) with exon hit	1.6
			Un 047560		1.6
		AW451454		adenylate kinase 3	
	321908	AA376936	Hs.20998	ESTS	1.6
	303349	AA382661		EST cluster (not in UniGene) with exon hit	1.6
25	324338	AL138357	Hs.247514		1.6
		AW300144		EST cluster (not in UniGene)	1.6
		A41000144			
	333193				1.6
	336433			CH22_FGENES.825_12	1.6
	312097	Al352096	Hs.157169	ESTs	1.6
30		AW204237		ESTs; Weakly similar to !!!! ALU SUBFAMILY J WARNING ENTRY !!!! [H.saplens]	
20					
		Al361722	Hs.192410		1.59
	308147	Al498991		EST singleton (not in UniGene) with exon hit	1.59
	313489	AA017492	Hs.135655	ESTs	1.59
	316289	AA902488	Hs.122952	ESTs ·	1.59
35		11002100	110.122002		1.59
JJ	326983	411000000		CH.21_hs gij5867657	
		AW205298	Hs.202372		1.59
	328397			CH.07_hs gi 5868397	1.59
	331970	AA461084	Hs.187677	ESTs	1.59
		N91419	Hs.12028		1.59
40					
40		Al292181	Hs.150036		1.59
	315921	Al147545	Hs.114172	ESTs	1.59
	322049	Al928242	Hs.144383	ESTs	1.59
		AA731518		EST cluster (not in UniGene) with exon hit	1.59
			Un 114000		1.59
15		Al026836	Hs.114689		
45	319142	F07366		EST duster (not in UniGene)	1.59
	313526	AW152263	Hs.249243	ESTs	1.59
	305937	AA883238		EST singleton (not in UniGene) with exon hit	1.58
	330123			CH.19_p2 gi 6671869	1.58
CO	327819			CH.05_hs gij5867968	1.58
50	318250	Al478814	Hs.134603	ESTs	1.58
	306760	AI034094	Hs.169476	tubulin; alpha; ubiquitous	1.58
	322358	AA220235	Hs.246836		1.58
	017000	Alconoco			1.58
	31/000	Al690269	Hs.201345		
	320725	AA703319	Hs.120967	ESTS	1.58
55	311332	AW292247	Hs.255052	ESTs	1.58
	334893			CH22 FGENES.452_7	1.58
		AA398215		EST duster (not in UniGene)	1.58
			11-004744		
		AW271639	Hs.221744		1.58
	303702	AW500748	Hs.224961	ESTs; Weakly similar to 73 kDA subunit of cleavage and polyadenylation	
60				specificity factor [H.sapiens]	1.57
	315086	Al492660	Hs.170935		1.57
		AA156499	Hs.8454	protein kinase; cAMP-dependent; regulatory; type II; alpha	1.57
	335549			CH22_FGENES.576_10	1.57
	329532			CH.10_p2_gi 3983505	1.57
65		AA180467		EST cluster (not in UniGene)	1.57
JJ			He 454500		
		Al801098	Hs.151500		1.57
	337896			CH22_EM:AC005500.GENSCAN.56-3	1.57
	330658	AA319514	Hs.211093	ESTs	1.57
		AI823969	Hs.132678		1.57

	317151	AW298195	Hs.255735	ESTs .	1.57
	308818	Al819700	Hs.208231	EST	1.57
	326547			CH.19_hs gi[5867307	1.57
_	318833	H06234	Hs.24888	ESTs	1.57
5		R31386		EST cluster (not in UniGene)	1.57
		Al124514		EST singleton (not in UniGene) with exon hit	1.57
	338083			CH22_EM:AC005500.GENSCAN.174-1	1.57
	316868	A1660898	Hs.195602	ESTs	1.57
4.0	310937	A1472880	Hs.170480	ESTs	1.57
10	328638			CH.07_hs gij6004473	1.57
		A1651039	Hs.148559	ESTs	1.56
	327058			CH.21_hs gi[6531965	1.56
		Al653733	Hs.204079	ESTs	1.56
		AF086529	•	EST cluster (not in UniGene)	1.56
15		A1745498	Hs.204579		1.56
		H49619	Hs.127301		1.56
	303841	A1934464		EST duster (not in UniGene) with exon hit	1.56
		AJ009849	Hs.199297	Homo sapiens GNAS1 gene encoding NESP55	1.56
00		AF156271		EST duster (not in UniGene)	1.56
20		AI052093	Hs.133132		1.56
		AL039604		EST cluster (not in UniGene) with exon hit	1.56
		AA833858		EST cluster (not in UniGene)	1.56
	328369			CH.07_hs gij5868388	1.56
05	329415			CH.Y_hs gi 5868874	1.56
25		AW468839	Hs.257767		1.56
	338794			CH22_EM:AC005500.GENSCAN.528-1	1.56
		AA243481	Hs.127320	ESTs; Weakly similar to KIAA0346 [H.sapiens]	1.56
		F08434		EST duster (not in UniGene)	1.56
20	334287	414400 4000	11. 000001	CH22_FGENES.369_17	1.56
30		AW024798	Hs.233374		1.55
		AA505833	Hs.162017		1.55
		AA682913	HS.24/1/9	ESTs; Weakly similar to KIAA0319 [H.sapiens]	1.55
		AA603092	11-040070	EST singleton (not in UniGene) with exon hit	1.55
35		AW502851	Hs.249978		1.55
33		AW501163	U= 04074	EST cluster (not in UniGene)	1.55 1.55
		H86709	Hs.21371	son of seventess (Drosophila) homolog 1	1.55
		AA761265	Hs.221281		1.55
		AA157392 Al299137	He 154014	EST cluster (not in UniGene) with exon hit	1.55
40	325389	MIZSSISI	Hs.154214		1.55
+0		AA417078	Hs.193767	CH.12_hs gi 5866921	1.55
		AA331732	Hs.137224		1.55
		AA258033	115.10/224	EST cluster (not in UniGene) with exon hit	1.55
		AA744875	Hs.189413		1.55
45		AA973297	Hs.126101		1.55
13		AI827065	Hs.224877		1.55
		T26438	113227077	EST singleton (not in UniGene) with exon hit	1.55
		AW160507		EST cluster (not in UniGene)	1.54
		AW138174	Hs.130651		1.54
50		AF086386	110.100001	EST cluster (not in UniGene)	1.54
-		AW411383	Hs.169688		1.54
	325303			CH.11_hs gi]5866908	1.54
		AI457663	Hs.128127		1.54
		AA582678		EST singleton (not in UniGene) with exon hit	1.54
55		AA861571		EST singleton (not in UniGene) with exon hit	1.54
		AA401367	Hs.128647		1.54
		Al381515	Hs.158381		1.54
		AA533505	Hs,185844		1.54
		AA513406	Hs.152307		1.54
60	323097	Z44354		guanine nucleotide binding protein (G protein); q polypeptide	1.54
	325043	W27919		Inositol polyphosphate-4-phosphatase; type I; 107kD	1.54
		Al376086	Hs.158759		1.54
	324573	AA491600	Hs.161942		1.54
		A1923673	Hs.212827		1.54
65		AA641092	Hs.257339		1.54
-		AF098363	2	EST cluster (not in UniGene) with exon hit	1.54
		Al459140	Hs.129109		1.54
	309536	AW151933		EST singleton (not in UniGene) with exon hit	1.54
•		AI146423	Hs.146709		1.53

	315674	AA651923	Hs.191850	ESTs ·	1.53
		N79341		EST cluster (not in UniGene)	1.53
		Al184510	Hs.143728	ESTs	1.53
,,,	330036				1.53
5		AA843868	Hs.190567		1.53
		AA972712	Hs.174818		1.53
		R51361	Hs.23423		1.53
		AA663591			1.53
10.	337685				1.53
10	335290	AIDEOCCZ			1.53
		Al858667 Al418246			1.53 1.53
		AW340374	He 121022		1.53
	335320	A11040074	113.121000		1.53
15	329841				1.53
		Al565071	Hs.159983		1.53
	332901				1.53
	305413	AA724659			1.53
	316707	AI016387	Hs.184406		1.53
20	313693	AW469180	Hs.170651		1.53
		AA922236	Hs.221037		1.53
		AF038966	Hs.184543		1.53
		Al248615	11- 450000		1.53
25		Al679968	Hs.152060		1.53
23		N27515 Al023175	Hs.40296	• •	1.53 1.53
	325958	AIU23175	Hs.167022		1.53
		AA664265	Hs.230213		1.53
		AW015667	Hs.119427		1.52
30		AA224368	Hs.185164		1.52
	301646	AA313954			1.52
	338752				1.52
		AW009312			1.52
25	301445	Ai208364	Hs.128233	ESTs; Weakly similar to REGULATOR OF CHROMOSOME	
35		******	11. 001450		1.52
		A1685263	Hs.201150		1.52 1.52
		AA635305 Al018150	Hs.121574 Hs.148781		1.52
	336205	A1010130	113.140701		1.52
40	325701				1.52
• •		AW189460	Hs.208358		1.52
	303121	AW407585	Hs.27769	ESTs; Weakly similar to mCAC [M.musculus]	1.52
	309271	Al986221		EST singleton (not in UniGene) with exon hit	1.52
4 ~	328385				1.52
45		Al318545			1.52
		AW103292	Hs.245328		1.52
		AA432067	Hs.258373		1.52 1.52
		AA232873 W52674		,,,,	1.52
50		AW298169	Hs.57553		1.52
-		AW207346	Hs.143202		1.52
		N63406	Hs.258697		1.52
		AF015950			1.52
·	314342	AJ873046	Hs.258775		1.51
55		AA887293			1.51
	301165	N85789	Hs.224155	ESTs; Weakly similar to PTERIN-4-ALPHA-CARBINOLAMINE	
	000040	A1000004	U- 040004		1.51
		AI932294	Hs.249004	ESTs; Weakly similar to B-CELL LYMPHOMA 6 PROTEIN [H.sapiens] ESTs; Weakly similar to SERINE/THREONINE-PROTEIN KINASE NRK2 [H.sapiens	1.51
60		Al554212 Al458207	Hs.174181		1.51
55		AL043148	Hs.186257		1.51
		AW139500	Hs.116135		1.51
		Al022056			1.51
	337976		•		1.51
65		A1083982			1.51
		A1569399	Hs.174746		1.51
		AA531082	Hs.240049		1.51
		AW025248	Hs.202445		1.51
	310759	AW135924	Hs.224883	rais	1.51

	310954	AW449044	Hs.171298	ESTs	1.51
	312019	T77046	Hs.188750	ESTs	1.51
	334773			CH22_FGENES.430_5	1.51
		AA490831	Hs.125056		1.51
5		AA296219		EST cluster (not in UniGene)	1.51
	337920			CH22_EM:AC005500.GENSCAN.67-3	1.51
	328993	•		CH.09_hs gi 5868536	1.51
	309245	AI972447		EST singleton (not in UniGene) with exon hit	1.51
	312172	Al222168	Hs.191168		1.51
10	304039	T47349		EST singleton (not in UniGene) with exon hit	1.5
	301329	Al149653	Hs.190496	ESTs	1.5
	313376	Al949246	Hs.200381	ESTs	1.5
	324248	AW504918		EST cluster (not in UniGene)	1.5
	308771	Al809301		EST singleton (not in UniGene) with exon hit	1.5
15	334935			CH22_FGENES.464_3	1.5
	319764	AA019827		EST cluster (not in UniGene)	1.5
	318519	T27135		EST cluster (not in UniGene)	1.5
	332807			CH22_FGENES.7_9	1.5
	322310	AF086376		EST cluster (not in UniGene)	1.5
20	324557	AA489166	Hs.156933		1.5
	332118	AA609585	Hs.162689		1.5
	319539			EST cluster (not in UniGene)	1.5
	313149	AW291092	Hs.201058		1.5
~-	329722			CH.14_p2 gi 6065785	1.5
25	323514			EST cluster (not in UniGene)	1.5
	308078	Al472621		EST singleton (not in UniGene) with exon hit	1.5
	337965			CH22_EM:AC005500.GENSCAN.100-10	1.5
	335905			CH22_FGENES.635_13	1.5

TABLE 14A shows the accession numbers for those primekeys lacking unigeneID's for Table 14. For each probeset we have listed the gene cluster number from which the oligonucleotides were designed. Gene clusters were compiled using sequences derived from Genbank ESTs and mRNAs. These sequences were clustered based on sequence similarity using Clustering and Alignment Tools (DoubleTwist, Oakland California). The Genbank accession numbers for sequences comprising each cluster are listed in the "Accession" column.

Pkey: CAT (Acces	iumber:	Unique Eos probeset identifier number Gene cluster number Genbank accession numbers					
Pkey	CAT number	Accession					
32206	4 234514_1	BE261397 Z78343 BE176419 AA383657 N90640 AA334052 AW955761 BE536232 ÆA374087 AA584776					
32140	9 197898_1	N71838 AA282003 T54072 AA761419 H92966 Al831371 Al095435 Al690247 R99331 AW964110 AA975590 AA346128 H94196 C03864					
32209	2 46678_1	AF085833 R69689 AW341677 AA923375 BE327566 AW630415 R69601 AW615339					
32145	2 212379_2	AW962489 H64300 AA329527					
31360	3 199797_1	AA284333 AW468119 AA284334 AA810992					
3208	6 36098_1	AB040928 T94673 Al289313 Al536039 Z44366 BE141499 D60116 D61488 D59945 AA419503 R28090 R72986 H03255 Al189112 Al912312 AW511018 Al401349 AW470144 C14624 Al335797 Z40300 Al014456 D60269 D60115 T16722 Al3706					
		D60270					
32213	9 46806 1	H53744 AF075088 H53797					
32150	0 552826_1	BE004271 Al248023 Al022157 H71999					
	3 441212_1	AA766346 AA809877 AA836116 AW469598 AW977404					
32221	5 47002_1	AF088005 N51816 N51731					
32223	5 47070_1	AF086106 AI193589 AW665594 N71795 AA722627 AW665373 AI300251					
	2 286374_1	AW812795 AA419617 H87827 AW299775 AW382168 AW382133 BE171659 AW392392 BE171641 AA541393					
31383	3 120893_1	AA766825 AA811180 AA085906 Al762946 AW977820					
	0 47376_1	AF086376 W77804 W72689 AA837735					
	3 47386_1	AF086386 W77947 W72708					
	2 47434_1	AF086431 AA886756 AI557237					
	1 47467_1	AF086467 W81444 W81445					
	5 47537_1	W95298 AF086529 Al912190 AW294159 Al458747 W94782					
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65		468554_1	AA830149 AW978407 M85983 AW503837 W00973 N56457 AW992226 T84921 R01342
0.0		1323199_1 1003489_1	R86913 R86901 H25352 R01370 H43764 AW044451 W21298
		1536467_1	F08434 Z42573 H26810
		765461_1	AI524124 R06841 R06842
		1534581_1	Z43108 F06295 R13085

	318885 94880_2 303841 79133_1	AA742999 Z43272 AA345258 AW956677 AA031942 W19657 BE616760 BE259848 BE382680 BE615587 AI934464 AA322745 T07155 AW961174 AA307302 Z41888 AA621992 AA188400 AW770608 AI147458 AI148408 AI696291 AA972591
5	303889 1777183_1 319539 63198_1 318905 1536408_1	T19204 T36109 T36107 R09027 AA344892 AA329574 AW955648 AW978708 Al567804 Al378935 AW014657 Al804134 R08922 N92947 BE546788 F08365 Z43395 R54298
	320187 396254_1 318996 65715 1	T99949 AA654769 AA664550 AW975264 Z44266 H06384 AV655948
10	319635 163534_1 319699 747196_1	R17531 AW960999 AA338366 AW673294 BE047729 BE047722 AA330746 AW841797 H05030 Al142105 R12654 Al458682 H24240 R14537 R18426 AW867082
	319713 1699356_1 319761 75324_2 319764 88596_1	R24204 R15712 T84695 AW630974 BE005208 R84237 AA724997 AA334867 AW955777 R18816 AA019827 R18947 H46852
15	319808 7069_3 321040 193331_1	T58960 AA609180 AA621130 Al927236 AA431075 AA261830 AW967855 H26953 AA262478
	320409 43709_1	AA226869 AA296516 AW959753 AA186390 AL359619 AA356195 AA148427 R22748 Al033624 BE548853 H95327 AW579751 BE561649 AA397533 BE617136 AA236444 T89946 AA247450 N55777 W38725 Al743846 Al808406 AA922229 Al051464 W04713 R11251 W19656 Al042319 AA489276 Al224533 H95274 AW269958 T89311 Al890088 Al862754
20		AI830968 AI669336 AI589780 AA534557 AW273839 AI338155 AI126632 N83542 BE046048 AA807028 AA848107 AW167978 AA976930 AA148428 AI289304 AI524262 AI625961 AA773469 AI222288 AI280054 AI242371 AA227222 AA973329 AA296517 AA829436 AA234526 AI149769 AI567865 AA936939 AI590681 AW469308 AI689531 AA486419
	319881 1585983_1 320488 368456 1	Al422051 Al057252 AA626941 Al475352 AW247913 Al222370 AA670122 AW198034 AA486418 Al363794 AA380739 H51299 H44619 H46391 R86024 H51892 T72744 Al817336 R32883 AA595590 Al743065 R31386
25	321121 1545647_1	W23285 H42714 F25381 F37215
	321205 81249_1	AA002047 N72537 H54142 H81580
	321253 375160_1 314043 155125_1	AA610649 Al699484 H59558 AA827082 AA732246 AA167611 AA830741
	320630 17685_2	AA199847 AA410224 R53323 AW936567 AW936569 AW936568 AW936571
30	313435 443527_1	AA769123 AA831715 AW977666 W92553
	313443 82292_1 313472 82811_1	AA005125 W95019 W93335 AA249037 AA007374 AA007466 Al816886
	321348 41762_1	Z49979 D61703 U30168
25	314138 179960_1	AA740616 AA654854 AA229923
35	320712 57156_2 321383 41924_1	R66867 R65678 R62673 W73128 R83101 AW968556 AJ238555 AW968731 AJ002574 AA459446 H70260 AW977557 AA767351 AW268572 AA810719 Al698677
	321303 41824_1	Al300460 AA907450 AA649224 T07415 Al536896 BE018515 Al279865 BE047421
	312996 187327_1	AW368634 Al702169 Al245179 AW368646 BE545574 AA249018 AW368633 N27553
40	306513 306537	AA989230 AA991705
40	306557	AA994530
	306598	AI000320
	306620	A1000929
45	306700 308078	A1022056 A1472621
73	306813	A1066544
	306830	AI075803
	306855	AI083982
50	329722 c14_p2 329728 c14_p2	
50	306890	A1092235
	308100	Al475949
	308147 306929	Al498991 Al124514
55	308352	A1610791
	308383	Al624497
	308521	A1689808
	308561 308617	AI701559 AI738720
60	308771	A1809301
-	308828	A1824829
	308896	A1858667
	303019 41850_1 303084 44211_1	AF098363 AF098365 AF174008 AF174027 AF174106
65	305092 AA642912	THE PERSON OF TH
	305169	AA663131
	305177	AA63591
	305235 305413	AA670480
	·	

	305849	AA861571
	305854	AA862733
	307113	Al183686
	307130	Al185234
5	305937	AA883238
•	305977	AA887293
	307451	Al248615
	307513	Al274307
	307848	Al364186
10	307871	Al368665
	307881	Al370434
	307932	AJ230822
	307944	Al418246
	307954	Al419692
15	307965	Al421641
	309245	AI972447
	309271	Al986221
	309365	AW072861
	309372	AW074330
20	309435	AW090537
	309506	AW137700
	309536	AW151933
	309709	AW242630
	325417 c12_hs	
25	325450 c12_hs	
	325452 c12_hs	
	309815	AW292760
	309839	AW296076
	309849	AW297444
30	309906	AW339340
	302705 31765_1	U09060 U09061
	304037 -	T26438
	304039	T47349
~~	304236	W93278
35	304257	AA053294
	304382	AA232873
	304405	AA282572
	304561	AA489792
40	304569	AA490934
40	304787	AA582678
	304921	AA603092
	327819 c_5_hs	
	304968	AA614308
4.5	306382	AA968967
45	331263 47479_1	AW780192 AA015718 W02571
	332252 1663967_1	N63882 T91174

5

TABLE 14B shows the genomic positioning for those primekeys lacking unigene ID's and accession numbers in Table 14. For each predicted exon, we have listed the genomic sequence source used for prediction. Nucleotide locations of each predicted exon are also listed.

10	Pkey: Ref: Strand:	Unique number corresponding to an Eos probeset Sequence source. The 7 digit numbers in this column are Genbank Identifier (GI) numbers Indicates DNA strand from which exons were predicted.
	Nt_position:	Indicates nucleotide positions of predicted exons.

				
15	Pkey	Ref	Strand	Nt_position
	332807	Dunham, I. et.al.	Plus	297686-297808
	332808	Dunham, I. et.al.	Plus	298277-298360
	332812	Dunham, I. et.al.	Plus	309688-310561
	332901	Dunham, I. et.al.	Pius	1841954-1842090
20	333149	Dunham, I. et.al.	Plus	3574317-3574413
	333916	Dunham, I. et.al.	Plus	8298994-8299169
	334026	Dunham, I. et.al.	Plus	9196549-9196681
	334061	Dunham, I. et.al.	Plus	9686941-9687077
	334073	Dunham, I. et.al.	Pius	9792201-9792374
25	334150	Dunham, I. et.al.	Plus	10529221-10529854
	334379	Dunham, I. et.al.	Plus	13908356-13908467
		Dunham, I. et.al.	Plus	15778859-15779026
	334773	Dunham, I. et.al.	Plus	16235169-16235328
20		Dunham, I. et.al.	Plus	19302753-19302881
30		Dunham, I. et.al.	Plus	20108247-20108373
		Dunham, I. et.al.	Plus	21491292-21491457
		Dunham, I. et.al.	Plus	22542132-22542246
		Dunham, I. et.al.	Plus	24935021-24935655
25		Dunham, I. et.al.	Plus	24990333-24990497
35		Dunham, I. et.al.	Plus	25044923-25045157
		Dunham, I. et.al.	Plus	29019796-29019877
		Dunham, I. et.al.	Plus	30051089-30051186 :
		Dunham, I. et.al.	Plus	31997555-31998040
40		Dunham, I. et.al.	Plus	23624127-23624224
40		Dunham, I. et.al. Dunham, I. et.al.	Plus Plus	32536159-32536395 2547461-3547246
		Dunham, I. etal.	Plus	3547161-3547245 3850500-3850643
		Dunham, I. et.al.	Plus	4113793-4113990
		Dunham, I. et.al.	Plus	7034267-7034392
45		Dunham, I. etal.	Plus	7166011-7166119
••		Dunham, I. et.al.	Plus	8072708-8072827
		Dunham, I. et.al.	Plus	10391398-10391600
		Dunham, I. et.al.	Plus	12205719-12205875
		Dunham, I. et.al.	Plus	12800037-12800181
50		Dunham, I. et.al.	Pius	19685043-19685354
	338506	Dunham, I. et.al.	Plus	21221871-21221953
	338794	Dunham, I. et.al.	Plus	27114697-27114763
	338910	Dunham, I. et.al.	Plus	28795375-28795551
		Dunham, I. et.al.	Plus	30760793-30760968
55	332864	Dunham, I. et.al.	Minus	1390386-1390296
		Dunham, I. et.al.	Minus	2035790-2035681
		Dunham, I. et.al.	Minus	3832993-3832494
		Dunham, I. et.al.	Minus	7286177-7286073
<i>(</i> 0		Dunham, I. et.al.	Minus	8523830-8523671
60		Dunham, I. et al.	Minus	8552629-8552330
		Dunham, I. et.al.	Minus	13294116-13293871
		Dunham, I. et.al.	Minus	13946021-13945781
		Dunham, I. et.al.	Minus	14432191-14432132
65		Dunham, I. et al.	Minus	19463909-19463815
UJ		Dunham, I. et.al. Dunham, I. et.al.	Minus Minus	21325792-21325667 21952922-21952826
	التعددن	omnani, L stat	willing	₹183€3€€-₹160€0€0

	335288	Dunham, I. et.al.	Minus	22304275-22303770
	335290	Dunham, I. et.al.	Minus	22309950-22309891
	335549		Minus	24666203-24666128
	335862		Minus	26690300-26690125
5	335864		Minus	26694537-26694382
	335905		Minus	26988888-26988719
	336205	•	Minus	30477456-30477311
	336276		Minus	32093320-32093181
10	336433		Minus	34067540-34067425
10	336605	Dunham, I. et.al.	Minus	15616509-15616358
	336616	Dunham, I. et.al.	Minus	26021027-26020848
	336679	Dunham, I. et.al.	Minus	2035790-2035681
	337043	Dunham, I. et.al.	Minus	17407330-17407251
	337272	Dunham, I. et.al.	Minus	28241476-28241307
15	337357	Dunham, I. et.al.	Minus	30906179-30906109
	337393	Dunham, I. et.al.	Minus	31471747-31471569
	337497	Dunham, I. et.al.	Minus	33371317-33371258
	337646		Minus	2648689-2648632
	337920		Minus	6051648-6051510
20		Dunham, I. et.al.	Minus	9318438-9318301
20		Dunham, I. et.al.	Minus	14166440-14166104
				26421374-26421135
	338752	Dunham, I. et.al.	Minus	
	338763		Minus	26628148-26628009
0.5	338983		Minus	29908865-29908702
25	339209	•	Minus	32492953-32492593
	325240		Minus	32301-32650
	329532		Pius	42937-43014
	329522	3983507	Minus	35265-35458
	329519	3983510	Plus	18407-18597
30	329511	3983514	Plus	20965-21325
	325326	5866875	Plus	47726-48024
	325303	5866908	Minus	73556-73630
	325389	5866921	Plus	239672-239759
	325417	5866925	Minus	110635-110745
35	325450	5866941	Minus	435379-435552
	325452	5866941	Minus	704103-704202
	325498		Plus	173372-173930
	325587	6682462	Plus	126724-126967
	325602	5866994	Plus	79122-79251
40	325701	5867028	Minus	72936-73046
	325780	6381953	Plus	63634-63873
	329722	6065785	Minus	112713-112992
	329728		Minus	207544-207741
	329666		Plus	98307-98446
45	329815		Minus	68431-68720
73	329841	6672062	Minus	40181-40331
			Minus	42450-42833
	325824			94358-94628
	325866		Minus	
50	325902		Minus	127729-127842
20	325958	5867142	Plus	53437-53550
	326014	••••	Minus	10358-10447
	329941		Minus	34319-34411
	330002		Plus	46097-46158
~ ~	326154		Minus	7103-7179
55		5867245	Pius	171799-171896
		5867269	Plus	75250-75903
	330036	6042048	Plus	117120-117216
	326547	5867307	Minus	623677-623870
	326495		Plus	11843-11930
60	326507	5867435	Minus	13038-13111
	326505	5867435	Minus	8818-8949
	326506	5867435	Minus	9368-9509
	326530	5867441	Minus	303000-303122
	326508		Plus	78904-79112
65		6671864	Minus	127553-127656
00		6671869	Minus	35311-35406
		6552462	Minus	69337-69670
		5867657	Minus	16023-16581
			Plus	1017630-1017788
	327014	2001004	rius	1017030-1017708

	326930	6456782	Plus	606950-607705
	326920		Minus	42425-42519
	327058		Plus	2384268-2384835
	327061		Minus	3486389-3486673
5.		6531965	Plus	4041318-4041431
J .		6531970	Minus	6-1088
•	330126		Plus	82458-82623
	327157		Minus	4408-4746
	327183		Plus	84317-84531
10			Minus	194652-194764
10	327192	5867445	Plus	48583-48773
	327288	5867481	Plus	145549-145708
		5867772	Minus	57796-58015
		6004459	Minus	97010-97123
15	327526		Plus	68767-69126
13	327574		Plus	141736-141900
		5867839	Plus	
	327752		Minus	93721-94421 92202-92717
		5867968	Pius	85267-85405
20		5867982		45203-45269
20		6671884	Plus	3982-4114
	330282		Plus	
		5868008	Plus	72807-72865
	328121		Plus	153782-153850 21082-21165
25		5868077	Plus	
23	328227		Minus	21082-21242
	327871		Minus	88889-89221
		5902482	Minus	542547-543133
	328624		Minus	120666-120836
30	328744		Plus	138639-138722
<i>5</i> 0	328799		Minus	80771-80923
	328291	5868363	Minus Plus	144244-144434
		5868375		191709-192239 75371-75583
-		5868388	Plus	369952-370155
35	328385		Plus	
33	328397	5868397	Plus	344967-345063
	328412		Plus	86427-86519
	328538		Plus	3814-4243
	328656		Plus Plus	792616-792729 294618-294903
40		6004473		
40	328903		Plus	23625-24468 38547-38837
	328960		Plus Minus	54458-54697
	330320	5932415 5868536	Plus	49160-50084
	329081		Plus	93368-93510
45		5868614	Plus	25805-26923
43			Plus	102168-102273
		5868626	Plus	166936-167020
	329192	5868716		71408-71707
		5868726	Minus	27422-27664
50		5868728	Plus	250541-250792
JU		5868732	Minus	
		5868874	Plus	1011438-1011818
	329454	5868887	Plus	51342-51593

TABLE 15: 169 GENES WITH SEQUENCE INFORMATION DEPICTED IN TABLE 16

Table 15 depicts UnigeneID, UnigeneTitle, Primekey, Predicted Cellular Localization, and Exemplar Accession for all of the sequences in Table 16. The information in Table 15 is linked by EosCode to Table 16.

Pkey: ExAcon:

Unique Eos probeset identifier number Exemplar Accession number, Genbank accession number

UnigenelD: Unigene number 10 Unigene Title: Unigene gene title EosCode: Internal Eos name

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Predicted cellular localization of gene product Localization:

15	Pkey	ExAccn	UnigeneID	Unigene Title	EosCode	Localization
	100394	D84276	Hs.66052	CD38 antigen (p45)	PBC1	plasma membrane
	100452	D87742	Hs.241552	KIAA0268 protein	PAB7	not determined
		L33881	Hs.1904	protein kinase C, iota	OAA1	cytoplasmic
20		M24736		selectin E (endothelial adhesion molecul	ACC5	plasma membrane
		M28214	Hs.123072	RAB3B, member RAS oncogene family	PFJ2	cytoplasmic
	101851	M94250	Hs.82045	midkine (neurite growth-promoting factor	LBH9	secreted
	102398	U42359		gb:Human N33 protein form 1 (N33) gene,	PDG3	
		U53347	Hs.183556	solute carrier family 1 (neutral amino a	PFJ4	plasma membrane
25	102669	U71207	Hs.29279	eyes absent (Drosophila) homolog 2	LEM9	cytoplasmic
	103119	X63629	Hs.2877	cadherin 3, type 1, P-cadherin (placenta	LBG2	plasma membrane
	103709	AA037316	Hs.13804	hypothetical protein d.1462O23.2	PD06	•
	104080	AA402971	Hs.57771	kallikrein 11	PBA6	secreted
	104144	AA447439	Hs.183390	hypothetical protein FLJ13590	PDM3	
30	104691	AA011176	Hs.37744	Homo sapiens beta-1 adrenergic receptor	PAV1	plasma membrane
	105370	AA236476	Hs.22791	transmembrane protein with EGF-like and	PDM9	plasma membrane
	106149	AA424881	Hs.256301	hypothetical protein MGC13170	PDO8	
	106579	AA456135	Hs.23023	ESTs	PAA4	plasma membrane
	107102	AA609723	Hs.30652	KIAA1344 protein	PAA3	not determined
35	107217	D51095		DKFZP586E1621 protein	PDG8	•
		AA054237		ESTs	PBF1	plasma membrane
	109014	AA156790	Hs.262036	ESTs, Weakly similar to Z223_HUMAN ZI	NC	PDG7
				hypothetical protein FLJ13782	BCU4	not determined
40		H04649	Hs.20843	Homo sapiens cDNA FLJ11245 fis, clone F		PDG4
40		H18836	Hs.31608	hypothetical protein FLJ20041	PAV9	plasma membrane
	-	T17185	Hs.83883	transmembrane, prostate androgen induce		CHA1 not determined
		T23855		KIAA1028 protein	PDO3	
		AA236545		cadherin-like protein VR20	PFJ6	plasma membrane
45		AA250737	Hs.72472	ESTs	BCY2	mitochondrial
45		AA599463		hypothetical protein MGC2648	PDV3	secreted
		AA609219		ESTs	OAB6	
		N41002	Hs.45107		PDT9	ER
		N51919	Hs.106778	ATPase, Ca++ transporting, type 2C, mem	D	PAJ5 not determined
50		N94303	HS.55028	ESTs, Weakly similar to I54374 gene NF2	PDM8	D400 - lanear
30		N95796		Homo sapiens prostein mRNA, complete c	US DDC0	-PAB2 plasma membrane
		R45175	Hs.117183		PBF8	_
		AA398246	Hs.97594		PDG5 PDV5	
		AA419011		prostate androgen-regulated transcript 1	BCU7	vesicular
55		AA428062	11- 00700	ESTs; protease inhibitor 15 (PI15)		PAZ1 not determined
22			Hs.98732		PD01	LWS1 Hot determined
				alpha-methylacyl-CoA racemase		PAA2 plasma membrane
			H- 000405	ESTs, Weakly similar to ALU1_HUMAN AL	DAVA	plasma membrane
		N62096	ns.293165	ESTs, Weakly similar to JC7328 amino aci		PDY4
60		AA128075 Al167942	Un 04020	transmembrane, prostate androgen induce	PAA5	plasma membrane
UU		R38438	Hs.61635	six transmembrane epithelial antigen of solute carrier family 15 (H+/peptide tra	PDO5	plasma membrane
			Hs.162859		PAA6	not determined
				secreted frizzled-related protein 4	BCX2	secreted
				calcium/calmodulin-dependent protein kin		2001010M
65		W26769	He 100700	CGI-86 protein	PAV6	vesicular
05		AA621604	113.103201	spondin 2, extracellular matrix protein	CJA5	not determined
	123303	ANUE 1004		Activities aving control income biorgin	June	nat domining

	129404	AA172056		ESTs	PAB4	
		R73640	Hs.11260	hypothetical protein FLJ11264	PAJ3	secreted
		AA128997	Hs.18953	phosphodiesterase 9A	PEE6	nuclear
_	131425	AA219134	Hs.26691	ESTs	PBA7	
5		AA031360		ESTs	PAA7	plasma membrane
		AA032221	Hs.61635	six transmembrane epithelial antigen of	PM17	plasma membrane
		U81599	Hs.66731	homeo box B13	PFJ5	nudear
		U42360	Hs.71119	Putative prostate cancer tumor suppresso		plasma membrane
10		X74331	Hs.74519	primase, polypeptide 2A (58kD)	PDM2	
10		U07919	Hs.75746	aldehyde dehydrogenase 1 family, membe		PDT1 mitochondrial
			Hs.75746	aldehyde dehydrogenase 1 family, membe		PDT1 mitochondrial
		AA045870		Homo sapiens mRNA; cDNA DKFZp564A0		PAB9 cytoplasmic
		U41060 AI800004	Hs.79136	LIV-1 protein, estrogen regulated	BCR4 PEU4	plasma membrane nudear
15		AI869666		hypothetical protein MAD (mothers against decapentaplegic, D		cytoplasmic
13		AA508353	He 105314	relaxin 1 (H1)	PBH3	secreted
			Hs 105887	ESTs, Weakly similar to Homolog of rat Z		500000
		D30891	Hs.19525	hypothetical protein FLJ22794	PBM4	not determined
		AW503733		KIAA1488 protein	PBY3	not determined
20	308050	A1460004	Hs.31608	hypothetical protein FLJ20041	PEU5	plasma membrane
	310382	A1734009	Hs.127699	KIAA1603 protein	PCQ8	
		A1420227		ESTs, Weakly similar to A46010 X-linked		plasma membrane _
			Hs.156142		PEN3	plasma membrane
25			Hs.140546		PCW3	
25		Al973051			PET5	
		AI682088		holocarboxylase synthetase (blotin-[prop	PBH8	
		AA861697 AI732100	Hs.120591 Hs.187619		PBY2 PBY1	•
			Hs.136319		BFF8	not determined
30	-	AV207200 AI538226		guanine nucleotide binding protein 4	CB07	cytopiasmic
50				ESTs, Weakly similar to TRHY_HUMAN Ti		PBM2not determined
		AW292425	110.222000	ESTs	PBM9	7 Dividilot dotominod
			Hs.134427		PBJ7	plasma membrane
	316442	AA760894	Hs.153023	ESTs	PBJ9	•
35	317548	Al654187	Hs.195704	ESTs	PBQ6	
				deoxyribonuclease II beta	PBQ7	
			Hs.159066	hypothetical protein FLJ10188	PBJ1	cytoplasmic
		AF071538		prostate epithelium-specific Ets transcr	PEN1	
40		AA460775		ESTs, Weakly similar to T17248 hypotheti		-1
40				ATP-binding cassette, sub-family C (CFTR	PEL9	plasma membrane plasma membrane
			3Hs.159330 Hs.31218		PBY4	not determined
				Homo sapiens LUCA-15 protein mRNA, sp		PBY8 not determined
		W07459	Hs.157601		CBF9	secreted
45				Homo sapiens cDNA FLJ12166 fis, clone it		PBQ1 not determined
			Hs.293616		PCQ7	plasma membrane
	323226	AF055019	Hs.21906	Homo sapiens clone 24670 mRNA sequen	œ	PCI2 not determined
	323287	AA639902	Hs.104215	ESTs, Moderately similar to SPCN_HUMA		PBJ5
50			Hs.143691		PBQ9	not determined
50				Homo sapiens cDNA: FLJ23241 fis, clone		PBY6 not determined
			Hs.292934		PBM3	ordania omia
			LI2' 192099	ESTs, Weakly similar to I38022 hypotheti gb:tt88f04.x1 NCI_CGAP_Pr28 Homo sapi		cytoplasmic PCW6
		Al685464 Al694767	He 120170	Homo sapiens cDNA FLJ13581 fis, clone i		PBJ4 plasma membrane
55		AI557019		small nuclear protein PRAC	CBK1	nuclear
	330211	7400.0.0	1101110	·	PBJ2	not determined
		U31382	Hs.299867	guanine nucleotide binding protein 4	PEW1	cytoplasmic
		AA449677	Hs.15251	hypothetical protein	PBM1	not determined
		T48536		TMPRSS2, transmembrane protease, seria	18	PEL3 plasma membrane
60	330892	AA149579	Hs.91202	ESTs	PBQ4	plasma membrane
		R36671	Hs.14846	Homo saplens mRNA; cDNA DKFZp564D0		PCQ1cytoplasmic
		N32912	Hs.291039		PCI4	nudear
		AA431407	Hs.98802	ESTs, Moderately similar to T14342 NSD1		not determined
65		N58172		gb:za21f09.s1 Soares fetal liver spleen	PBQ5	nuclear RR III act determined
OD		AA340504		gb:hw31a09.x1 NCI_CGAP_Kid11 Homo s		PBJ8 not determined
	332798	T94885		transgelin 2	PBQ8 PBH2	secreted nuclear
	334447		•	•	PBY9	not determined
	338255				PBY7	not determined
	J-1-2-00			•		

	401424				PFG2	mitochondrial
	407122	H20276	Hs.31742	ESTs	PEW7	
		S79876	Hs.44926	dipeptidylpeptidase IV (CD26, adenosine	PEZ3	plasma membrane
		AF216077		Homo sapiens clone HB-2 mRNA sequence		PEY1
5		AK000631		hypothetical protein FLJ20624	PFG1	nuclear
•		NM 00598		sine oculis homeobox (Drosophila) homolo		nuclear
		U80034	Hs.68583	mitochondrial intermediate peptidase	PEZ9	mitochondrial
		BE244589		glyoxalase I	PFJ3	cytoplasmic
			Hs.246973		OBH6	Cytopidaniic
10				Homeo box A13	PFC6	
10			Hs.130853		PEZ5	
		X57010	Hs.81343	"collagen, type II, alpha 1 (primary ost	PFJ1	secreted
		AA279490		calmegin	-	.ER
		AJ820961			PFA1	.EN
15		NM_00499			PEY4	
15				ATP-binding cassette, sub-family C (CFTR		
		U24577	Hs.93304	"phospholipase A2, group VII (platelet-a	PFH9	secreted
	42100/	AVV 10145U	HS.109201	CGI-86 protein	PFH2	plasma membrane
				*arachidonate 15-lipoxygenase, second ty		cytoplasmic
20		AW102723		guanylate cyclase 1, soluble, alpha 3	PFA3	
20			9Hs.154424	"delodinase, lodothyronine, type II"	PFH6	secreted
		AF030880		solute carrier family, member 4	PFD4	plasma membrane
		AA418000		potassium intermediate/small conductance		plasma membrane
				KIAA0575 gene product	PFD6	nudear
25		AA460421		ESTs	PEZ7	•
25			Hs.119383		PEY5	
		BE245562		adrenergic, beta-2-, receptor, surface	PEZ4	plasma membrane
				Rho GTPase activating protein 6	PFG6	nuclear
		D89053			PEZ1	
20		NM_00274	2Hs.2891	protein kinase C, mu	PFH4	cytoplasmic
30		AA527941		gb:nh30c04.s1 NCI_CGAP_Pr3 Homo sap		PFA2
		AI669973	Hs.200574		PEW8	
		W07088	Hs.293685		PFG3	
			Hs.325198		PEY3	
25		A1446444			PEW5	
35		AI972867	Hs.7130	copine IV	PEW6	
		AA909358	Hs.128612		PFC8	
		AB028945		cortactin SH3 domain-binding protein	PEZ6	
		AF126245	Hs.14791	"acyl-Coenzyme A dehydrogenase family, o		PFH7
40		AF035269			PFH8	
40				prostate cancer antigen 3, non-coding DD		
		NM_014253		odz (odd Oz/ten-m, Drosophila) homolog 1		plasma membrane
		AF055575	Hs.23838		PFD2	plasma membrane
	451939		Hs.27311	single-minded (Drosophila) homolog 2	PFJ8	
45	451982		Hs.27373	Homo sapiens mRNA; cDNA DKFZp56401		PFG9plasma membrane
45		AI922988			PFD8	
		NM_002202		ISL1 transcription factor, LIM/homeodoma		nuclear
					PFC5	cytoplasmic
	452946	X95425	Hs.31092	EphA5	PFH3	plasma membrane

TABLE 15A shows the accession numbers for those primekeys lacking a unigeneID in Table 15. For each probeset we have listed the gene cluster number from which the oligonucleotides were designed. Gene clusters were compiled using sequences derived from Genbank ESTs and mRNAs. These sequences were clustered based on sequence similarity using Clustering and Alignment Tools (DoubleTwist, Oakland California). The Genbank accession numbers for sequences comprising each cluster are listed in the "Accession" column.

1	Λ
1	v

Pkey:	Unique Eos probeset Identifier number
CAT number:	Gene cluster number
Accession:	Genbank accession numbers

15

15			
	Pkey	CAT number	Accession
20	116393	131543_1	Al972402 Al634409 Al523716 Al799749 W44518 Al424438 Al688513 Al971048 Al686324 AW013854 AA588483 AA528111 Al627428 Al582200 Al669296 Al826926 Al620526 Al669958 Al972458 Al924500 AA512903 W44517 AA335363 AW238997 BE300165 BE250665 AA284195 AA523420 W52834 Al471970 Al952824 AW003820 AW009463 AA669796 AA114966 Al653342 AA115038 Al342150 Al092100 Al968211 W51994 Al804005 Al201420 Al123210 Al738405 Al674964 Al970341 AW027500 Al493316 Al333193 Al139353 AA599463 Al656163 Al804200 Al365321 Al990213 Al657011 AA650025 Al968810 Al341978 AA599839 AW592602 AA644289 Al468578 Al565265 Al565228 BE221535 AW973052
25	101485	18113_1	AA296520 AL021940 M30640 NM_000450 M24736 M61894 AL047443 H39560 Al694691 AA916787 Al214796 AA939085 Al150616 AA412553 AA412545 Al051015 T27654 AA694430
30	126399	17331_1	AA088767 AF224278 AA128075 AL035541 AA027926 AI761441 AI972096 AW071693 AI742327 AI377498 AI804815 AI640802 AI885001 AI921394 AA595115 N71820 AI921217 AW007283 AI467828 AI369306 AA917446 AI493698 AA088701 AA126899 AI936228 AW204238 AI039567 AI925027 BE138909 AW452945 AW135998 AA310984 AA027860 AW073519 AI537597 AA953976 AI521341 AW273569 AW050740 AA536113 AA559064 AI474392 AW135709 AA535181 AW572959 AA570597 AI905464 AI677810 AI587642
30	400004		AW975102 AA424310 AA482527 N64192 AA658276 AW889117 AA486591 AW889172 AI381990 AI381991 AI673419 AI990950 AA487031 AI272934 AI150565 AA229168 AW316722 AI142707 BE222396 AA614168 AA122026 AW338227 AA632457 AI968726 AW369662 AA512956 AA541675 AA451748 AI250993 BE146418 AA122025
35		94346_1 21074_1	AI362575 AI805082 AW263421 AI432462 AA135870 AA031360 AA031604 AA298475 AA298464 NM_012445 AB027466 BE407510 BE047605 AA047125 AW084003 AA149494 AA149490 AA292528 AA570505 AA526186 AW006250 AW007762 AI341557 AI799666 AI972710 AI377966 AI962810 AI084783 AI458032 AI190971 AW148913 AA372354 AW970032 AW007426 AA650188 AI123203 AI122890 AI280975 W73595 W73495 AI863238 AA374109 AA603986 AW149089 AW957523 AI307748 AI921067 AI336463 F24537 AI380460 AI367500 AI189309 AI814701 AI766921 AW572106 AA037024 AW0772576 AA578293 AI288103 AA235464 AW450642 AA574230 AW294024 AI589229 AI580733 AW512227 AA877009 AI660255 AW188597 AA558228
40			ALS72782 AA658397 AL274628 AI866359 AA864573 AI264439 AA621604 AW515493 AW243333 Z39737 AI567038 AA573997 AA573559 AW236431 AI652870 AI884973 AA034505 AA047126
45		156454_1 9836_1	Al267700 Al720344 AA191424 Al023543 Al469633 AA172056 AW958465 AA172236 AW953397 AA355086 AL080235 AA031750 D81382 Al480231 Al095947 Al560953 BE010721 Al670290 AA374945 AA125792 D51527 D51556 Al685541 D51559 AW117286 AA195741 Al675138 AW593439 Al201885 T30590 AW952100 D51095 AA523864 W70043 AA987586 Al421515 Al205532 AA127069 Al337367 D51595 Al453785 AW075677 AW088359 C14287 C14287 C14284
		19266_1	AF163474 NM_016590 AF163475 AI761105 AI770098 AA410580 AA411616 AI590343 AI739050 AL050198 AI862645 AA419104 AA513809 AA333032 AI816915 AW139625 AA640889 AI311391 AI627693 AW135514 AA419011 AI269149 AI245259 AI970008 AI970017 AW139445 AA569503 AI761072 AI766179 AI759995 AI300776 AI870129 AW150770 AA226501 AA226220
50		291015_1 entrez_U42359	Al249368 Al742316 AA428062 AA442089 Al864189 BE349478 Al803475 Al584049 BE552085 Al088609 Al264197 Al886144 Al129474 Al307145 BE181300 AW058403 Al696838 AW748598 AA442196 Al216428
	315051 324626	347217_1 336411_1	AW292425 BE467167 AI702953 BE550961 BE222309 AI299348 AI693336 AA541708 AI685464 AW971336 AA513587 AA525142
55	319191	16065_1	NM_012391 AF071538 AB031549 Al685592 Al745526 AA662204 AW130657 AA662164 AW971121 Al668916 AA513274 Al991223 Al979170 AW298436 AA639821 Al859010 AW513942 Al687669 AA662521 AA548598 Al345056 Al305374 BE043418 Al432856 Al334840 Al379796 Al492693 Al307915 BE042082 Al307834 Al307858 Al309488 BE042210 Al435670 Al371605 Al862491 Al284563 Al306872 Al255044 Al254601 Al251236 Al473073 Al473042 Al432760 Al435664 Al336826 Al289365 Al369096 Al862274 Al334871 Al349863 Al250405 Al377617 Al309895 Al313017 Al862291 Al311936 Al378718 Al305722 Al306769 Al308888 Al334565 Al862296
60			AI344230 AI435685 AI344087 AI378696 AI311209 AI435775 AI310611 AI311154 AI432289 AI431561 AI492681 AI432867 AI335288 AI492796 AI432769 AI310299 AI432273 AI379820 AI275319 AI435753 AI609441 AI432767 AI369100 AI311420 AI349974 AI247157 AI334677 AI270910 AI224320 AI305608 AI334489 AI377152 AI350012 AI370086 AI335053 AI306781 AI306750 AI334849 AI334874 AI340380 AI307876 AI305974 AI305972 AI311521 AI334872 AI862509 AI311498 AI335051 AI289684 AI310859 AI311862 AI862483 AI492775 AI307906 AI492708 AI289693 AI340373 AI307910 AI311359 AI435653 AI334865 AI311492 AI492809 AI492690 AI431576
65			Al862268 Al311879 Al308435 Al492792 Al862512 Al275321 Al431568 Al431564 Al307885 Al307926 Al435692 Al435778 Al310182 Al308894 Al492707 Al492713 Al308560 Al307829 Al343234 Al580598 AW472796 Al340918 Al310243 Al309368 Al307920 Al289665

			· · · · · · · · · · · · · · · · · · ·
			Al306777 AW086318 AW086292 AW086378 Al310027 Al275293 Al369082 Al340900 Al306749 Al371558 AW086287 BE043803 Al306793 Al306272 Al287948 Al270917 Al284816 Al336813 Al284546 Al308044 Al275290 Al270872 Al306795 Al289687 Al223570
			AISO5303 AIS06272 AIS0746 AIS7647 AIS64816 AIS36813 AIS64946 AIS0644 AIS76260 AIS7672 AIS0672 AIS06795 AIS65607 AIS6567 AIS657 AIS67 AIS657 AIS67 A
_			Al305357 Al275270 Al345932 Al436549 Al307925 Al311502 Al344238 Al343182 Al308508 Al305988 Al270790 Al379792 Al305647
5		,	Ai305410 Ai432251 Ai436517 Ai343227 Ai305534 Ai340387 Ai271043 Ai305499 Ai271046 Ai305962 Ai289465 Ai305378 Ai289725
			AI310848 AI305848 AI289362 AI252964 AI307049 AI310831 AI306993 AI306796 AI224659 AI305969 AI349855 AI306164 AI306948
			A1284676 A1309155 A1343202 A1432785 A1306815 A1369081 A1270885 A1289699 A1435704 A1309647 A1305716 A1311281 A1287927 A1472995 A1340423 A1270958 A1307069 A1305364 A1270807 A1275306 A1311890 A1275263 A1432750 A1289371 A1432861 A1255113
			AI305709 AI473008 AI311168 AI309711 AI377164 AI271201 AI289560 AI309710 AI306195 AI311201 AI287741 AI271066 AI432876
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			Al473160 Al432903 Al223720 Al254979 Al334862 Al306926 Al269541 Al432248 Al435722 Al435698 Al432859 Al310683 Al473175 Al335144 Al289467 Al436489 Al306928 Al473033 Al305763 Al307868 Al307882 Al348959 Al435736 Al432857 Al432896 Al435735
			AISS3144 RE29407 AIAS0408 AIS0320 AIATS055 AIS07056 AIS07068 AIS07062 AIST36959 AIATS750 AIAS2690 AIATS755 AIATS765 AIAT
15			Al432279 Al432882 Al334670 Al436512 Al432827 Al432852 Al473051 Al473077 Al435697 Al271509 Al492781 Al472983 Al473018
			Al432897 Al473043 Al432871 Al436536 Al473157 Al349715 Al432777 Al473016 Al473158 Al340369 Al307941 Al432773 Al377146
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			Al289550 Al305721 Al340870 Al270901 Al308575 Al307904 Al340715 Al270941 Al309808 Al246867 Al473014 Al307039 Al289360 Al473069 Al492786 Al344013 Al305876 Al436510 Al340742 Al473028 Al307891 BE041871 BE041268 BE042340 BE041946
20			BE041783 Al306173 Al201948 Al926972 Al275769
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		C_5_p2	6 5 LINK C4G1.G
			O_S_CINN_C4GT.G G_387_7_LINK_EM
25		372969_1	AA669097 AA513815 AA026798 AA676526 AA704429 AA704269 AW118292 AA579216 N58172
	332396	20265_1	AW579842 BE156562 BE156690 BE156489 BE081033 AK001559 BE149402 M85387 AW367811 AW367798 R17370 Al908947
			AA382932 R58449 H18732 AA371231 AW962899 AA713530 AW892946 R53463 H11063 AW068542 Z40761 BE176212 BE176155 W23952 W92188 AW374883 AA303497 AW954769 AA036808 BE168063 AW382073 AW382085 AL041475 H80748 A078161
			BE463983 AI805213 AI761264 W94885 N94502 AI623772 AI419532 AI810302 AI634190 AW002516 AW150777 AI352312 AI367474
30			AW204807 Al675502 Al337026 AW134715 BE328451 Al123157 Al560020 Al300745 Al608631 Al248873 AA742484 AW051635
			H18646 Al245045 AA507111 Al640510 Al925594 AA115747 AA143035 AA151106
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			AW895564 AW895597 AW895595 AW895665 AW888518 AI903724 F06081 F08503 AL119462 AW895730 AW888516 R26511
35			R26489 AA334126 AA327626 N85713 AW895998 AA223622 F05468 AA370749 W05590 M78202 AA371073 AW498607 R15017
			T16991 AA001282 AA001138 AA551566 AA330159 AI922855 AA383512 AA029603 D82246 D82171 T94933 H56545 AA348060
			AA176888 R96764 AW451817 AA385766 AA452618 AI690057 AA988822 BE549928 AA150901 W57992 AW899925 C05281 AA932042 AA370980 AW962877 W04741 AA369982 AW385948 AA922466 N75882 AI422070 AI361256 AI680224 D57122 T94885
			R53266 R46713 T19071 AW796277 AA325333 F04719 F02334 AA358146 AA626597 AA358304 AW028099 AL119570 D57290
40			D58273 D57796 N48555 Al361969 AA329457 D57225 AW024046 AA992606 AW022118 AW021538 AA935845 H89870 H56546
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			AA493436 Al365636 BE464751 AW149384 AA102442 AW771368 Al818251 Al126368 D51049 Al421542 Al559467 AW079779
			AW021048 AW023969 AW044214 Al458264 AA027274 Al620254 AW028917 BE219511 AA326242 N67561 Al971273 AA878328
45			D57131 AA770662 Al309299 Al796767 AA613338 W58076 Al566287 Al445573 Al880260 AA001919 AW339259 Al492610 Al492611
			R97692 Al301425 AA722603 D58361 Al350323 AA973926 Al431263 AA516126 AA865467 Al925177 N39443 AA001943 Al299371 Al082412 AA665090 AA583433 H89871 AA977231 Al362219 Al056096 Al270446 N67524 N22103 AW614224 AA744054 AW243622
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~ 0			AI034036 AI582196 AW264534 AI418961 AA570761 AI343538 AA650341 AA992503 AA770004 AL039666 AI862675 AW190335
50			AA610274 AW418627 BE467472 D56786 T28749 Al217610 Al359556 T23523 AL040189 AA846222 AA651636 D51280 Al888986
			AI521167 AI340177 AW612815 AI625285 AA621607 AA177059 AA229768 AA829768 AI749682 AW190631 N75299 AA230089 AI915632 BE069542 AA890020 AA528397 AA995390 BE503860 AA570812 AW339396 AI197986 AI203725 AI282379 AA670375
			AA461513 F01728 AW243599 C00856 N75567 R95995 AA150932 R95991 AA648060 AA933800 AA927073 AA101126 AA864190
			T93566 BE167472
55		25529_1	AF030880 NM_000441 AC002467 AA385554 H23053 AW891838 AI139968 AA653057 AI695233
		342819_1 6391 1	AA527941 Al810608 Al620190 AA635266 AB028945 T77648 F13328 AL157605 Z46212 AA304736 F11855 T66098 T30174 AW954164 AW176301 AW748243 AA456428
	4-0464	0331_1	Al369958 AA938565 AW959613 Z42008 AA994779 Al683909 F11019 F10926 Al769597 Al752550 T65015 Al884314 AA643954
			Z41838 AW020147 Al038822 AW571822 AA299781 AA894928 AF131790 BE005411 Al902476 AW082695 AA464384 R42750
60	447040	7110 4	AW902301 AA464273 R05837 Z38294 H41098 AL134507 M86079
	44/210	7119_1	AF035269 AF035268 NM_015900 T96213 U37591 AA156832 AA299371 A1084325 H95977 AI765967 BE221465 AA156726 AI969563 AW024539 AI436791 AI949451 AA843093 AI452756 AA824232 AI306667 T96131 AW207447 AW243556 AW957032 AI084332
			H95978 U30998
<i></i>	449625	8113_1	NM_014253 AF100772 BE088769 AL022718 BE161779 AW863569 BE161640 AL039060 BE168542 AW296554 AA323193 AA235370
65			AW779760 N48674 AI375997 R45432 D59344 AI203107 F07491 R35360 R25094 AI913631 AI498402 T61382 AI016320 N45526
	452020	89513_1	T61415 AA331486 . AI922988 H05475 AA021608 AW169947 AA913750 Z41614 AW800012
	702000		**************************************

TABLE 15B shows the genomic positioning for those primekeys lacking unigene ID's and accession numbers in Table 15. For each predicted exon, we have listed the genomic sequence source used for prediction. Nucleotide locations of each predicted exon are also listed.

10	Pkey: Ref: Strand: Nt_posi	Se pul sec Ind	quence source. blication entitled quence of huma licates DNA stra	rresponding to an Eos probeset The 7 digit numbers in this column are Genbank Identifier (GI) numbers. "Dunham I. et al." refers to the "The DNA n chromosome 22." Dunham I. et al., Nature (1999) 402:489-495. Ind from which exons were predicted. e positions of predicted exons.
15	Pkey	Ref	Strand	Nt_position
	334447	Dunham, i. et.al.	Pius	14308764-14308824
	332798	Dunham, I. et.al.	Minus	232147-231974
	338255	Dunham, I. et.al.	Minus	15242294-15242231
$\Delta \Delta$				

59158-59215

24223-24428

Plus

Plus ·

5

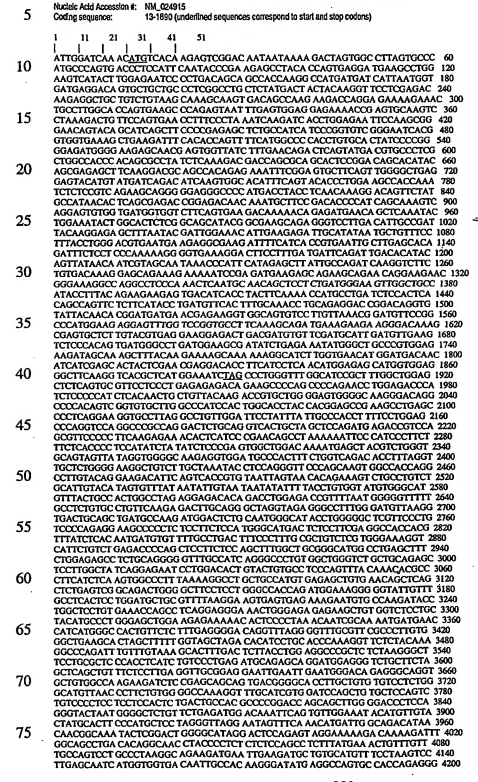
20

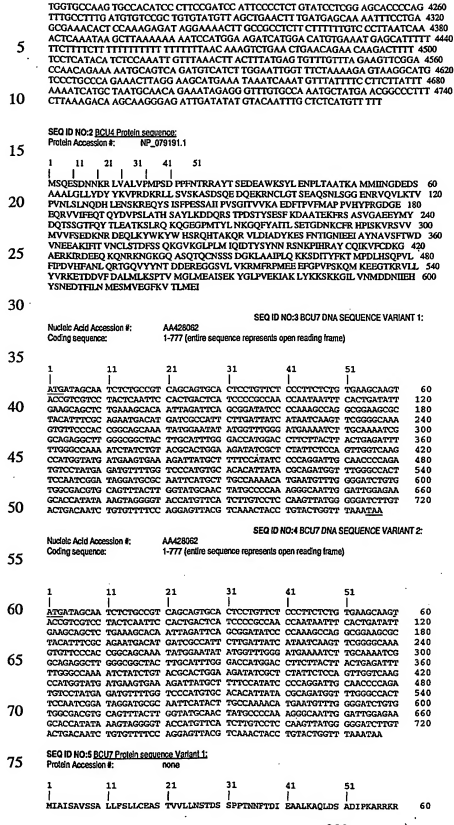
330211 6013592

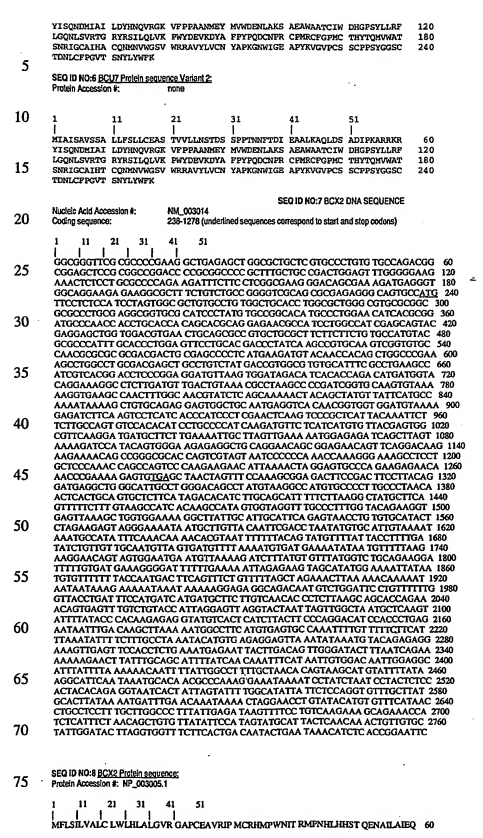
401424 8176894

TABLE 11 AND SEQUENCE LISTING

SEQ ID NO:1 BCU4 DNA SEQUENCE







5 KKKTAGRTSR SNPPKPKGKP PAPKPASPKK NIKTRSAQKR TNPKRV SEQ ID NO:9 CBK1 DNA SEQUENCE Nucleic Acid Accession #: NM_032391 10 Coding sequence: 129-302 (underlined sequences correspond to start and stop codons) 51 11 21 31 15 GTCCTTCCTC TCCTAGCCTA AGGCGTGCAA ACAGAGCGCC ACTGGGAGGC TGAAACCTTT 60 AGGCCGATGC TTGCTTGCAA GGTCAGGCAA GCTGGATTCT GGTCCCCACC TTTGCAGAGA GAACAGCGAT GTTGTGCGCC CATTTCTCAG ATCAAGGACC GGCCCATCTT ACTACCTCCA 180 AGAGTGCTTT TCTCTCTAAT AAGAAAACAT CTACTTTGAA ACATCTACTG GGCGAGACCA 240 GGAGTGATGG CTCAGCCTGT AATTCTGGAA TTTCGGGAGG CCGAGGCAGG AAGATTCCTT 300 20 GAGCACAGGA GTTCCAGACC AGCCTGGGCA ATGTAGCAAG ACGCTGTCTC TATTTATACA 360 ATAAAATTTT TTTAAAAAAG G 25 SEQ ID NO:10 CBK1 Protein sequence: Protein Accession #: NP_115767 21 41 51 11 31 30 MLCAHFSDQG PAHLTTSKSA FLSNKKTSTL KHLLGETRSD GSACNSGISG GRGRKIP SEQ ID NO:11 CHA1 DNA SEQUENCE Nucleic Acid Accession #: NM_020182 35 Coding sequence: 96-854 (underlined sequences correspond to start and stop codons) 40 TCCTTGGGTT CGGGTGAAAG CGCCTGGGGG TTCGTGGCCA TGATCCCCGA GCTGCTGGAG 60 AACTGAAGGC GGACAGTCTC CTGCGAAACC AGGCA<u>ATG</u>GC GGAGCTGGAG TTTGTTCAGA 120 TCATCATCAT CGTGGTGGTG ATGATGGTGA TGGTGGTGGT GATCACGTGC CTGCTGAGCC 180 ACTACAAGCT GTCTGCACGG TCCTTCATCA GCCGGCACAG CCAGGGGCGG AGGAGAGAAG ATGCCCTGTC CTCAGAAGGA TGCCTGTGGC CCTCGGAGAG CACAGTGTCA GGCAACGGAA 300 45 TCCCAGAGCC GCAGGTCTAC GCCCGCCTC GGCCCACCGA CCGCCTGGCC GTGCCGCCCT 360 TOGOCCAGOG GGAGOGOTTO CACOGOTTOC AGCCCACOTA TOCGTACOTG CAGCACGAGA 420 TCGACCTGCC ACCCACCATC TCGCTGTCAG ACGGGGAGGA GCCCCCACCC TACCAGGGCC 480 CCTGCACCCT CCAGCTTCGG GACCCCGAGC AGCAGCTGGA ACTGAACCGG GAGTCGGTGC 540 GCGCACCCC AAACAGAACC ATCTTCGACA GTGACCTGAT GGATAGTGCC AGGCTGGGCG 600 50 GCCCCTGCCC CCCCAGCAGT AACTCGGGCA TCAGCGCCAC GTGCTACGGC AGCGGCGGGC 660 GCATGGAGGG GCCGCCGCCC ACCTACAGCG AGGTCATCGG CCACTACCCG GGGTCCTCCT 720 TCCAGCACCA GCAGAGCAGT GGGCCGCCCT CCTTGCTGGA GGGGACCCGG CTCCACCACA 780 CACACATCGC GCCCCTAGAG AGCGCAGCCA TCTGGAGCAA AGAGAAGGAT AAACAGAAAG 840 GACACCCTCT CTAGGGTCCC CAGGGGGGCC GGGCTGGGGC TGCGTAGGTG AAAAGGCAGA 900 55 ACACTCCGCG CTTCTTAGAA GAGGAGTGAG AGGAAGGCGG GGGGCGCAGC AACGCATCGT 960 GTGGCCCTCC CCTCCCACCT CCCTGTGTAT AAATATTTAC ATGTGATGTC TGGTCTGAAT GCACAAGCTA AGAAGACTTG CAAAAAAAAA AAGAAAAAAG AAAAAAAAA ACCACGTTTC 1020 1080 TTTGTTGAGC TGTGTCTTGA AGGCAAAAGA AAAAAAATTT CTACAGTAAA AAAAAAAAA 60 SEQ ID NO:12 CHA1 Protein sequence: Protein Accession #: NP 064567 65 21 31 41 51 MAELEFVQII IIVVVMMVMV VVITCLLSHY KLSARSFISR HSQGRRREDA LSSEGCLWPS 60 ESTVSGNGIP EPQVYAPPRP TDRLAVPPFA QRERFHRFQP TYPYLQHEID LPPTISLSDG 120 EEPPPYQGPC TLQLRDPEQQ LELNRESVRA PPNRTIFDSD LMDSARLGGP CPPSSNSGIS 180 70 ATCYGSGGRM EGPPPTYSEV IGHYPGSSFQ HQQSSGPPSL LEGTRLHHTH IAPLESAAIW SKEKDKQKGH PL SEQ ID NO:13 CJA5 DNA SEQUENCE 75 Nucleic Acid Accession #: NM_012445 276-1271 (underlined sequences correspond to start and stop codons) Coding sequence: 31 51 11 80 1 305

YEELVDVNCS AVLRFFFCAM YAPICTLEFL HDPIKPCKSV CQRARDDCEP LMKMYNHSWP 120 ESLACDELPV YDRGVCISPE AIVITOLPEDV KWIDITPDMM VQERPLDVDC KRLSPDRCKC 180 KKVKPTLATY LSKNYSYVIH AKIKAVQRSG CNEVTTVVDV KEIFKSSSPI PRTQVPLITN 240 SSCQCPHILP HQDVLIMCYE WRSRMMLLEN CLVEKWRDQL SKRSIQWEER LQEQRRTVQD 300

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       LVRLRQSPRA FIPPAPVLPS RDNEIVDSAS VPETPLDCEV SLWSSWGLCG GHCGRLGTKS
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50
                         26-457 (underlined sequences correspond to start and stop codons)
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SEQ ID NO:17 LEM9 DNA SEQUENCE

1-1617 (underlined sequences correspond to start and stop codons) Coding sequence: 5 21 31 51 11 ATGGTAGAAC TAGTGATCTC ACCCAGCCTC ACTGTAAACA GCGATTGTCT GGATAAACTG 60 AAGITTAACC GTGCTGACGC TGCTGTGTGG ACTCTGAGTG ACAGACAAGG CATCACCAAA 120 10 TCGGCCCCC TGAGAGTGTC CCAGCTCTTC TCCAGATCTT GCCCACGTGT CCTCCCCCGC 180 CAGCCTTCCA CAGCCATGGC AGCCTACGGC CAGACGCAGT ACAGTGCGGG GATCCAGCAG 240 GCTACCCCCT ATACAGCTTA CCCACCTCCA GCACAAGCCT ATGGAATCCC TTCCTACAGC 300 ATCAAGACAG AAGACAGCTT GAACCATTCC CCTGGCCAGA GTGGATTCCT CAGCTATGGC 360 TCCAGCTTCA GCACCTCACC CACTGGACAG AGCCCATACA CCTACCAGAT GCACGGCACA 420 15 ACAGGGTTCT ATCAAGGAGG AAATGGACTG GGCAACGCAG CCGGTTTCGG GAGTGTGCAC 480 CAGGACTATC CTTCCTACCC CGGCTTCCCC CAGAGCCAGT ACCCCCAGTA TTACGGCTCA 540 TCCTACAACC CTCCCTACGT CCCGGCCAGC AGCATCTGCC CTTCGCCCCT CTCCACGTCC 600 ACCTACGTCC TCCAGGAGGC ATCTCACAAC GTCCCCAACC AGAGTTCCGA GTCACTTGCT 660 GGTGAATACA ACACACAA TGGACCTTCC ACACCAGCGA AAGAGGGAGA CACAGACAGG 20 CCGCACCGGG CCTCCGACGG GAAGCTCCGA GGCCGGTCTA AGAGGAGCAG TGACCCGTCC 780 840 CCGGCAGGGG ACAATGAGAT TGAGCGTGTG TTCGTGTGGG ACTTGGATGA GACAATAATT ATTITICACT CCTTACTCAC GGGGACATTT GCATCCAGAT ACGGGAAGGA CACCACGACG 900 TCCGTGCGCA TTGGCCTTAT GATGGAAGAG ATGATCTTCA ACCTTGCAGA TACACATCTG 960 TTCTTCAATG ACCTGGAGGA TTGTGACCAG ATCCACGTTG ATGACGTCTC ATCAGATGAC 1020 25 ANTICCCIAG ATTTAGCAC ATACAACTTC TCCGCTGACG GCTTCCACAG TTCGGCCCCA 1080 GGAGCCAACC TGTGCCTGGG CTCTGGCGTG CACGGCGGCG TGGACTGGAT GAGGAAGCTG 1140 GCCTTCCGCT ACCGCCGGT GAAGGAGATG TACAATACCT ACAAGAACAA CGTTGGTGGG 1200 1260 ACAGACCTCT GGCTGACCCA CTCCCTGAAG GCACTAAACC TCATCAACTC CCGGCCCAAC 1320 30 TGTGTCAATG TGCTGGTCAC CACCACTCAA CTAATTCCTG CCCTGGCCAA AGTCCTGCTA 1380 TATGGCCTGG GGTCTGTGTT TCCTATTGAG AACATCTACA GTGCAACCAA GACAGGGAAG 1440 1500 GAGAGCTGCT TCGAGAGGAT AATGCAGAGA TTCGGCAGAA AAGCTGTCTA CGTGGTGATC GGTGATGGTG TGGAAGAGGA GCAAGGAGCG AAAAAGCACA ACATGCCTTT CTGGCGGATA 1560 TCCTGCCACG CAGACCTGGA GGCACTGAGG CACGCCCTGG AACTGGAGTA TTTA<u>TAG</u> 35 SEQ ID NO:18 LEM9 Protein sequence: NP 005235 Protein Accession #: 40 41 51 11 21 31 MVELVISPSL TVNSDCLDKL KFNRADAAVW TLSDROGITK SAPLRVSQLF SRSCPRVLPR 60 QPSTAMAAYG QTQYSAGIQQ ATPYTAYPPP AQAYGIPSYS IKTEDSLNHS PGQSGFLSYG SSPSTSPTGQ SPYTYOMHOT TGFYQGGNGL GNAAGFGSVH QDYPSYPGFP QSQYPQYYGS SYNPPYVPAS SICPSPLSTS TYVLQEASHN VPNQSSESLA GEYNTHNGPS TPAKEGDTDR 180 45 240 PHRASDGKLR GRSKRSSDPS PAGDNEIERV FVWDLDETII IFHSLLTGTF ASRYGKDTTT 300 SVRIGLMMEE MIFNLADTHL FFNDLEDCDQ IHVDDVSSDD NGQDLSTYNF SADGFHSSAP 360 420 GANLCLGSGV HGGVDWMRKL APRYRRVKEM YNTYKNNVGG LIGTPKRETW LQLRAELEAL TDLWLTHSLK ALNLINSRPN CVNVLVTTTQ LIPALAKVLL YGLGSVFPIE NIYSATKTGK 50 ESCFERIMOR FGRKAVYVVI GDGVEEEQGA KKHNMPFWRI SCHADLEALR HALELEYL SEQ ID NO:19 OAA1 DNA SEQUENCE Nucleic Acid Accession #: NM_002740 55 178-1968 (underlined sequences correspond to start and stop codons) Coding sequence: 31 51 60 CCGCGGTTCC GGCTGCTCCG GCGAGGCGAC CCTTGGGTCG GCGCTGCGGG CGAGGTGGGC AGGTAGGTGG GCGGACGGCC GCGGTTCTCC GGCAAGCGCA GGCGGCGGAG TCCCCCACGG CGCCCGAAGC GCCCCCGCA CCCCCGGCCT CCAGCGTTGA GGCGGGGGAG TGAGGAGATG 180 CCGACCCAGA GGGACAGCAG CACCATGTCC CACACGGTCG CAGGCGGCGG CAGCGGGGAC 240 CATTCCCACC AGGTCCGGGT GAAAGCCTAC TACCGCGGGG ATATCATGAT AACACATTTT 300 65 GAACCTTCCA TCTCCTTTGA GGGCCTTTGC AATGAGGTTC GAGACATGTG TTCTTTTGAC 360 420 AACGAACAGC TCTTCACCAT GAAATGGATA GATGAGGAAG GAGACCCGTG TACAGTATCA TCTCAGTTGG AGTTAGAAGA AGCCTTTAGA CTTTATGAGC TAAACAAGGA TTCTGAACTC 480 TTGATTCATG TGTTCCCTTG TGTACCAGAA CGTCCTGGGA TGCCTTGTCC AGGAGAAGAT 540 AAATCCATCT ACCGTAGAGG TGCACGCCGC TGGAGAAAGC TTTATTGTGC CAATGGCCAC 70 ACTITICCAAG CCAAGCGTTT CAACAGGCGT GCTCACTGTG CCATCTGCAC AGACCGAATA TGGGGACTTG GACGCCAAGG ATATAAGTGC ATCAACTGCA AACTCTTGGT TCATAAGAAG 720 TGCCATAAAC TCGTCACAAT TGAATGTGGG CGGCATTCTT TGCCACAGGA ACCAGTGATG 780 CCCATGGATC AGTCATCCAT GCATTCTGAC CATGCACAGA CAGTAATTCC ATATAATCCT 840 TCAAGTCATG AGAGTTTGGA TCAAGTTGGT GAAGAAAAG AGGCAATGAA CACCAGGGAA 900 75 AGTGGCAAAG CTTCATCCAG TCTAGGTCTT CAGGATTTTG ATTTGCTCCG GGTAATAGGA 960 AGAGGAAGTT ATGCCAAAGT ACTGTTGGTT CGATTAAAAA AAACAGATCG TATTTATGCA 1020 ATGAAAGTTG TGAAAAAAGA GCTTGTTAAT GATGATGAGG ATATTGATTG GGTACAGACA 1080 GAGAAGCATG TGTTTGAGCA GGCATCCAAT CATCCTTTCC TTGTTGGGCT GCATTCTTGC 1140 TITCAGACAG AAAGCAGATT GITCTITGTT ATAGAGTATG TAAATGGAGG AGACCTAATG 1200 80 TTTCATATGC AGCGACAAAG AAAACTTCCT GAAGAACATG CCAGATTTTA CTCTGCAGAA 1260

Nucleic Acid Accession #: NM_005244

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25	WIDEEGDPCT RRWRKLYCAN CGRHSLPQEP	VSSQLELEEA GHTFQAKRFN VMPMDQSSMH	21 AYYRGDIMIT FRLYELNKDS RRAHCAICTD SDHAQTVIPY	ELLIHVFPCV RIWGLGRQGY NPSSHESLDQ	PERPGMPCPG KCINCKLLVH VGEEKEAMNT	EDKSTYRRGA KKCHKLVTIE RESGKASSSL	60 120 180 240
30	SNHPFLVGLH HERGI IYRDL YGFSVDWWAL ASVLKSFLNK	SCFQTESRLF KLDNVLLDSE GVLMFEMMAG DPKERLGCHP	LVRLKKTDRI FVIEYVNGGD GHIKLTDYGM RSPFDIVGSS QTGFADIQGH	LMFHMQRQRK CKEGLRPGDT DNPDQNTEDY PFFRNVDWDM	LPEEHARFYS TSTFCGTPNY LFQVILEKQI MEQKQVVPPF	AEISLALNYL IAPEILRGED RIPRSLSVKA	300 360 420 480 540
35	DNFDSQFTNE	PVQLTPDDDD	IVRKIDQSEF			DNA SEQUENCE	
40	Nucleic Acid Acc Coding sequence	ession #: L05628 :: 197-47) 92 (underlined sec	ruences correspon	d to start and stop	codons)	
40							
40	1	11	21	31	41	51	
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45	CCAGGCGGGG GCCGCCGGG TGCCGCGGC CGCCCGGGCC CTGGGACTGG CACGGTCCTC CTATCTCTCC TGCCTTGGGA AAGAAGTCGG CACCACGCTG	TTGCGGCCCC CCGCCGCCCC CGCCGCGCC ACCGGCATGG AATGTCACGT GTGTGGGTGC CGACATGACC TTTTTGCTGT GGCATATTCC CTTGCTACCT	GGCCCGGCT GGCTAGCGCC GGCTCCGGGG GGAATACCAG CTTGTTTTTA GAGGCTACAT GGATCGTCT TTGCCCCAGT TTTTAATTCA	CCTGCGCCG AGCAGCCGGG CCGATCACC CTTCTGCAGC CAACCCCGAC CCTCTGGGCC TCAGATGACA CTGGGCAGAC GTTTCTGGTC GCTGGAGAGG	CCGCCGCCGC CCCGATCACC CGCCGCCCGG GCCGATGGCT TTCACCAAGT TGTTTCCCCT CCTCTCAACA CTCTTCTACT AGCCCAACTC AGGAAGGAG	51 CGCCGCCGC CGCCGCCGG TGCCCGCCG CCGACCGGT GCTTTCAGAA TCTACTTCCT AAACCAAAAC CTTTCTGGGA TCTTGGGCAT TCTAGTCTTC	120 180 240 300 360 420 480 540
45 50	CAGGCGGGG GCCGCCGCGC CGCCCGCGC CGCCCGCGC CTGGGACTGG CACGGTCCTC CTATCTCTCC TGCCTTGGA AAGAAGTCGG AAGAAGTCGG AAGAAGTCGG	TTGCGGCCCC CCGCCGCCAC CCGCCGCGCC ACCGCCATGG AATGTCACGT GTGTGGGTGC CGACATGACC TTTTTGCTGT GGCATATTCC CTTCGTACCT CTCACTTTCT	GGCCCCGGCT GGCTAGCGCC AGCAACCGGG CGCTCCGGGG GGAATACCAG CTTGTTTTTA GAGGCTACAT GGATCGTCTG TGGCCCAGT TTTTAAATTCA GGCTGGTAGC	CCCTGCGCCG AGCAGCCGGG CCCGATCACC CTTCTGCAGC CAACCCCGAC CCTCTGGGCC TCAGATGACA CTGGGCAGAC GTTTCTGGTC CCTGGAGAGG CCTAGTGTGTT	CCGCCGCCGC CCCGATCACC CCCCGCCCGG GCCGATGGCT TTCACCAAGT TGTTTCCCCT CCTCTCAACA CTCTTCTACT AGCCCAACTA AGGAAGGAG GCCCTAGCCA	51 CGCCGCCCC CGCCGCCCGC TGCCCGCCC CCGACCCGC TCTACTTCCT AAACCAAAAC CTTTCTGGGA TCTTGGCAT TCCATCTCTC TCCCTGAGATC	120 180 240 300 360 420 480 540 600 660
45	CAAGCGGCG GCCGCCGCG GCCGCCGCC CGCCCGCGC CTGGGACTGG CATCTTCTCC TGCCTTGGGA AAGAAGTCGG CACCACGCTG AGGGATCATG CAAAATTATG	TTGCGGCCCC CCGCCGCCAC CCGCCGCGCC ACCGCATGG AATGTCACGT GTGTTGGGTGC CGACATGACC TTTTTGCTGT GCGATATTCC CTTGCTACCT CTCACTTTTCT ACAGCCTTAA	GGCCCGGCT GGCTAGCGCC GGCTCCGGGG GGAATACCAG CTTGTTTTTA GAGGCTACAT GGATCGTCT TTGCCCCAGT TTTTAATTCA	CCTGCGCCG AGCAGCCGGG CCGATCACC CTTCTGCAGC CAACCCCGAC CTCTGGGCAC CTGGGCAGAC GTTTCTGGTC GCTGGTGTGT CCTAGTGTGT CCAGGTGGAC	CGGCGCGCGC CCCGATCACC CGCCGCCGG GCCGATGGCT TTCACCAAGT TGTTTCCCCT CCTCTCAACA CTCTTCTACT AGCCAACTC AGGAAGGAA GCCCTAGCCA CTGTTTCGTG	51 CGCCGCCGC GCCGCCGC GCCGCCGC CCGACCGCT GCTTTCAGAA TCTACTTCCT AAACCAAAAC CTTTCTGGGA TCTTGGGCAT TCCAGTCTTC TCCTGAGATCT ACATCACTTT	120 180 240 300 360 420 480 540
45 50	CAGGCGGGG GCCGCCGCGC CGCCCGCGCC CGCCCGCGCC CTGGGACTGG CACGGTCCTC TGCCTTGGGA AAGAAGTCGG CACCACGCTG AGGGATCATG CAAAATTATG CAAAATTATAC	TTGCGGCCC CCGCCGCAG CCCCGCGCC ACCGCATGG AATGTCACGT GTGTGGGTGC CGACATGACC TTTTTGCTGT GGCATATTCC CTTGCTACCT CTCACTTTAC ACAGCCTTAA ACTGCCTCT TCGGAAACCA	GGCCCGGCT GGCTACCGGG GCTCCGGGG GGAATACCAG GGTCCTACTTTTA GAGGCTACAT GGATCGTCTG TGGCCCAGT TTTTAATTCA GGCTGGTAGC AAGAGGATGC TACTCATTCA TCCACGACCC	CCCTGCGCCG CCCGATCACC CTTCTGCAGC CAACCCCGAC CCTCTGGGCC TCAGATGACA CTGGGCAGAC GTTTCTGGTC GCTGGAGAGG CCTAGTGTGT CCAGGTGGAC GCTCGTCTTG TAATCCCTGC	CCGCCGCCGC CCCGATCACC CCCGCCCGG GCCGATGGCT TTCACCAAGT TGTTTCCCT CCTCTCAACA CTCTTCTACT AGGAAGGAG GCCCTAGCCA CTGTTTCGTG TCCTGTTTCTC CCAGAGTCCA	51 CGCCGCCGC CGCCGCCGC CGCCGCCGG CCGACCGGC CCGACCGGC CCTTCCAAAA CCTACCTCCT AAACCAAAAAC CTTTCTGGGA TCTTGGGCAT TCAGTCTTC CCTGACATC ACATCACTTT CAGATCGCCC CAGACTCCCTT CAGATCGCCCT CGCCTTCCTT	120 180 240 300 360 420 480 540 600 660 720 780 840
45 50	CAAGCGGCG GCCGCCGCG CGCCGCGCC CGCCGCGCC CTGGGACTGG CATCTCTCC TGCCTTGGA AAGAAGTCGG CACCACGCTG AGGGATCATG CAAAATTATG CTACGTCTAC ACCCCTGTTAC	TTGCGGCCCC CCGCCGCAG CCGCCGCAG ACCGGCATGG AATGTCACGT CGACATGACC TTTTTGCTGT GGCATATTCC CTTGCTACCT CTCACTTTTCT ACAGCCTTAA TTTTCCTCT TCGGAAACCA ATCACCTTCT	GGCCCGGCT GGCACCGGG GCAACCGGG GGAATACCAG CTTGTTTTTA GAGGCTACAT GGATCGTCTT TTTTAATTCA GGCTGGTAGC AAGAGGATGC TACTCATTCA TCCACGACCC GGTGGATCAC	CCCTGCGCCG AGCAGCCGGG CCCGATCACC CTTCTGCAGC CAACCCGAC CCTCTGGGCA CTGGGCAGAC CTTGGTGTG CCTAGAGAGG CCTAGTGTGT CCAGGTGGAC GCTCGTCTTG CAGGTTGAC AGGGTTGAT AGGGTTGAT AGGGTTGAT AGGGTTGAT AGGGTTGAT AGGGTTGAT AGGGTTGAT AGGGTTGAT	CGGCGCGCGC CCGATCACC CGCCGCCGG GCCGATGGCT TTCACCAAGT TGTTTCCCCT CCTCTCAACA CTCTTCTACT AGCCAACTC AGGAGGGAG GCCTAGCCA CTGTTTCGTG TCCAGAGTCCA GTCCGGGGCT	51 CGCCGCCGC GCCGCCGG TGCCCGCG CCGACCGCT GCTTTCAGAA TCTACTTCCT AAACCAAAAC CTTTCTGGGA TCTTGGGCAT TCCTGAGATC TCCTGAGATC ACATCACTTT CAGATCGCTC GCGCTTCCTT ACCGCCAGCC	120 180 240 300 360 420 480 540 660 720 780 840 900
45 50 55	CCAGGCGGCG GCGCCGCCGCGCGCGCGCGCGCCGCGCC	TTGCGGCCCC CCGCCGCCAG CCGCCGCGCC ACCGCCATGG AATGTCACGT GTGTTGGTTGC TTTTTGCTGT GCGATATTCC CTTGCTACCT CTCACTTTTCT ACAGCCTTAA TTTTCCTCT TCGGAAACCA ATCACCTTCT AGTGACCTTCT AGTGACCCTTCT	GGCCCGGCT GGCACCGGG CGCTCCGGG GGAATACCAG GGATCGTTTTA GGAGCTACAT GGATCGTCT TGTCACTC TGCCCAGT TTTTAATTCA GGCTGGTAGC AAGAGGATGC TACTCATTCA TCCACGACCG GGTGGATCAC GGTGGTACAC GGTGGTACAC	CCCTGCGCCG AGCAGCCGGG CCCGATCACC CTTCTGCAGC CAACCCGAC CTGGGCAGAC CTGGGCAGAC GTTTCTGGTC CCTAGTGTGT CCAGGTGGAC GCTCGTCTTG AGGGTTGAT CCAGGTTGAC CAGGTTGAC CAGGTTGAC CAGGTTGAC CAGGTTGAC CAGGTTGAC CAAGGAGGAC CAAGGAGGAC	CGGCGCGCGC CCCGATCACC CGCCGCCGG GCCGATGGCT TTCACCAAGT TGTTTCCCCT CCTCTCAACA CTCTTCTACT AGCCAACTC AGGAAGGAA GCCTAGCCA CTGTTTCGTG TCCTGTTTCT CCAGAGTCCA ACGCGGGCCT ACGTCGGAAC	51 CGCCGCCGC GCCGCCGC GCCGCCGC CCGACCGCC GCTTCAGAA TCTACTTCCT AAACCAAAAC CTTTCTGGGA TCTAGGTCTT TCAGACTCT TCCTGAGATC ACATCACTT CAGATCGCC ACATCACTCT ACGCCTCCCT ACGCCAGCC AAGTCGTGCC	120 180 240 300 420 480 540 600 660 720 780 840 900 960
45 50	CAGGCGGGG GCCGCCGCGC CGCCCGCGCC CGCCCGCGCC CTGGGACTGG CACGGTCCTC TGCCTTGGGA AAGAAGTCGG CACCACGCTG AGGGATCATG CAAAATTATG CTACGTCTAC ACCCCTGTTC CCTGCAGGG CCTGGAGGGC TGTTTTGGTA TGTTTACTAC	TTGCGGCCC CCGCGCCAC CCGCCGCGCC ACCGCATGG AATGTCACGT TTTTGCTGT GGCATATTCC CTTGCTACCT CTTACTTTCT ACAGCCTTAA ATTTCCCTCT TCGGAAACCA ATCACCTCT AGAACTGAC TCCAAGGATC	GGCCCGGCT GGCTAGCGC AGCAACCGGG GGCTCCGGGG GGAATACCAG GGTCGTACAT GGATCGTCTG TGGCCCAGT TTTTAATTCA GGCTGGTAGC AAGAGGATGC TACTCATTCA TCCACGACCC GGTGGATCAC GGTCCTTAAA AGAAGGATGC CTGCCCAGCC CTGCCCAGC	CCCTGCGCCG CCCGATCACC CTTCTGCAGC CAACCCCGAC CTCTGGGCC TCAGATGACA CTGGGCAGAC GTTTCTGGTC GCTGGAGAGG CCTAGTGTGT CCAGGTGGAC GCTCGTCTTG TAATCCCTGC AGGGTTGAT CAAGGAGGAC CGCCAAGACT GAAAGAGAGT	CCGCCGCCGC CCCGATCACC CCCCGCCCGG GCCGATGGCT TTCACCAAGT TGTTTCCCT CCTCTCAACA CTCTTCTACT AGGAAGGAG GCCCTAGCCA CTGTTTCGTG TCCTGTTTCT CCAGAGTCCA GTCCGGAGC TCCTCGGAGC TCCAGGGGCT ACGTCGGAAGCAC TCCAAGGTGG	51 CGCCGCCCCC CGCCGCCCGC TGCCCGCCGC CCGACCCGCT CCTTTCAGAA TCTACTTCCT AAACCAAAAC CTTTCTGGCAA TCTTGGCAAT TCCTGAGATC TCCTGAGATC ACATCACTTT CAGATCGCTC GCGCTTCCTT ACCGCCAGCC AAGTCGTCCT ACGCCAGCC AAGTCGTCAC AGTCGTAAGGT ATGCGAATGA	120 180 240 300 360 420 480 540 660 720 780 840 900 960 1020
45 50 55	CAGGCGGGG GCCGCCGCGC CGCCGCGCC CGCCCGCGC CTGGGACTGG CACGTCCTC CTATCTCTCC TGCCTTGGGA AAGAATCGG CACCACGCTG AGGGATCATG CAAATTATG CAAAATTATG CAACCCTGTTC CCTCTCGAGG CCTGGAGGGC TGTTTTTGGTA TGTGACTCC GGAGGTGGAG	TTGCGGCCCC CCGCCGCAG CCGCCGCAG ACCGCATGG AATGTCACGT GTGTGGGTGC CCACATGACC TTTTTGCTGT GCCATATTCC CTTGCTACCT ACAGCCTTAA TTTTCCTCT TCGGAAACCA ATCACCTTCT AGTGAACCTCT AAGAACTGGT ACCAGGAT TCCAAGGAT CCCAAGGAT GCTTTGATCC GCTTTGATCCT TCGGAAACTGT AGTGAACTTCT AGTGAACTGT AGTGACCTTCT AGTGACCTTCT AGTGACCTTCT AGTGACCTCT AGTGACCTCT AGTGACCTCT AGTGACCTCT GCTTTGATCG GCTTTGATCG	GGCCCGGCT GGCACCGGG GGAATACCAG GGTCCGGGG GGAATACCAG CTTGTTTTTA GAGGCTACAT TGTAATTCA GGCCCAGT TACTCATCA AGAAGGATGC CAGGCC GGTGGATCAC GGTCGTAAT CCACGACC GGTGGATCAC GGTCCTTAAA AGAAGGATCC TCCCCAGCC TCAAGTCCCC	CCCTGCGCCG CCGACCCGGC CCACCCGAC CTTCTGCAGC CCTCTGGGCC CTCTGGGCAC CTGGGCAGAC CTGGGCAGAC CCTAGTGTGT CCAGGTGGAC AGGGTTGATT CAAGGAGGAC CGCCAAGACT CAAAGAGAGAC ACAGAAGACA ACAGAAGAGACA ACAGAAGAGAA	CGGCGCGCGC CCGATCACC CGCCGCCGG GCCGATGGCT TTCACCAAGT TGTTTCCCCT CCTCTCAACA CTCTTCTACT AGCCAACTC AGGAAGGAG GCCTAGCCA CTGTTTCGTG TCCTGTTTCT CCAGAGTCCA GTCCGGGCT ACGTCGGAAC AGGAAGCAC TCCAAGGTGG TCCAAGGTGG TCCAAGGTGG TCCAAGGTGG	51 CGCCGCCGC GCCGCCGG GCTCCGCG GCTTCAGAA TCTACTTCCT AAACCAAAAC CTTTCTGGGA TCTAGTCTT TCCAGACTTT TCAGTCTT CAGACTCTT ACACTCTT ACACTCTT ACACTCCT ACACTCCTT ACCGCCAGCC AAGTCGTCCC GGGTTAAGGT ATGCGAATGA CTCTGTTTAA	120 180 240 300 360 420 480 540 660 720 780 840 900 960 1020 1080 1140
45 50 55 60	CAGGCGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGG	TTGCGGCCCC CCGCCCAG CCGCCAGGC ACCGCATGG AATGTCACGT CGACATGACC TTTTTGCTGT GGCATATTCC CTCGCTACT ACAGCCTTAA TTTTCCTCT TCGGAAACCA ATCACCTTCT AGGAACCT AAGAACTGGA TCCAAGGATC CCTCAAGGATC AAGAACTGGA ACCACTTTG AAGAACTGGA TCCAAGGATC AAGAACTGGA AAGACTGGA AAGACTTTG AAGACCTTTT	GGCCCGGCT GGCATACCGG GGAATACCAG GGATCGTTTTTA GAGGCTACAT GGATCGTCT TGCCCAGT TTTTAATTCA GGCTGGTAGC AAGAGGATGC TACTCATTCA TCCACGACCC GGTGGTACAC GGTCCTTAAA AGAAGGATGC TCAGGACCC TCAGGTCCC GGCCTACTT	CCCTGCGCCG AGCAGCCGGG CCCGATCACC CTTCTGCAGC CAACCCGAC CTTCTGGGCAGAC CTTGGGCAGAC GTTTCTGGTC CCTAGTGTGT CCAGGTGGAC GCTCGTCTTG CAGGTGGAC GCTCGTCTTG CAGGTGGAC GCTCGTCTTG CAAGGAGGAC CGCCAAGACT CAAGAAGGAGAC CCTCATGAGAGGAC CCTCATGAGAGGAC CCTCATGAGC	CGGCGCGCGC CCCGATCACC CGCCGCCGG GCCGATGGCT TTCACCAAGT TGTTTCCCCT CCTCTCAACA CTCTTCTACT AGCCAACTC AGGAAGGAC CTGTTTCGTG TCCTGTTTCT CCAGAGTTCG AGGAAGCAAC AGGAAGCAAC TCCAAGGTGG TCCAAGGTGG TCCAAGGTGG TCCAAGGTGG TGGAACCCCT	51 CGCCGCCGC CGCCGCCGC CGCCGCCGC CCGACCGCCT GCTTTCAGAA TCTACTTCCT AAACCAAAAC CTTTCTGGGA TCTAGGTCTTC CAGATCACTCTT CAGATCGCC CGCTTCCTC CGCGTTCCTC CGGCTTCCTC ACACCCC AAGTCGTCC CGGTGAAGGT ATGCGAATGA ACTCGTATAA AGGCCATCCA	120 180 240 360 420 480 540 660 720 780 900 960 1020 1080 1140 1200
45 50 55	CAGGCGGGGGGGGGGGGGGGGGGTGGAGGGGTGGAGGGGGGGG	TTGCGGCCCC CCGCCCAG CCGCCAG ACCGCATG AATGTCACGT TTTTTGCTGT GGCATATCC CTTGCTACCT ACAGCCTTAA TTTTCCTCT ACAGCCTTAA ATCACCTTCT AGGAAACCA ATCACCTTCT AAGAACTGGA TCCCAAGGATC CCTTTGGTACCTT AAGAACTGA ACCACTTTTTTCCAAGGATCC AAGAACTGA AAGACCTTTG AAGAACTTTG AAGAACTTTG AAGAACTTTG AAGAACTTTG AAGAACTTTG AAGACCTTTG AAGAACCTTTG AAGAACTTTG AAGACCTTTG AAGACCTTTG AAGACCTTTG AAGACCTTTG AAGACCTTTG AAGACCTTTCC	GGCCCGGCT GGCTAGCGG GGATACCAG GGATACCAG GGATCGTTTTA GGATCGTCTG TGGCCCAGT TTTTAATTCA TCGCTGGTAGC AAGAGGATGC TACTCATTCA TCCACGACCC GGTGGATCA GGTCCTTAAA AGAAGGATG CTGCCAGCC TCAAGTCCC GGCCCAGCT CCACGACCC GGCCCACT CCACGACCC CCAGGACCC GGCCCTACTT CGCCCAGCC TCAAGTCCCC GGCCCTACTT GGCCCAGAT	CCCTGCGCCG AGCAGCCGGG CCCGATCACC CTTCTGCAGC CAACCCGAC CTTCTGGGCC CTTGGAGAGACG CCTGGAGAGGC CCTGGTGTT CCAGGTGGAC GCTGGTGTGT CCAGGTGAC GCTCGTCTTG CAGGAGGAC AGGGTTGATT CAAGGAGGAC CGCCAAGACT GAAAGAGGGC CCTCATGAGC CCTCATGAGC CCTCATGAGC CTTAAAGTTG CCTCATGAGC CTTAAAGTTG	CCGCCGCCGC CCCGATCACC CCCCATCACC CCCCATCACT TTCACCAAGT TGTTTCCACC CTCTTCTACT AGCCAACTC AGGAAGGAG GCCTAGCCA CTGTTTCTT TCCTGTTTCT CCAGAGTCCA GTCCGGGGCT ACGTCGGAAC AGGAAGCAGC TCCAAGGTGG TCCAAGGTGG TCCAAGGTGG TCCAAGGTGG TCCAAGGTGG TCCAAGGTGG	51 CGCCGCCGC GCCGCCGG GCTCCGCG GCTTCAGAA TCTACTTCCT AAACCAAAAC CTTTCTGGGA TCTAGTCTT TCCAGACTTT TCAGTCTT CAGACTCTT ACACTCTT ACACTCTT ACACTCCT ACACTCCTT ACCGCCAGCC AAGTCGTCCC GGGTTAAGGT ATGCGAATGA CTCTGTTTAA	120 180 240 300 360 420 480 660 660 720 840 900 91020 1020 1140 1200 1260 1320
45 50 55 60	CAGAGGGGGGGGGGCGCGCGCGCGCGCGCGCGCGCGCCGC	TTGCGGCCCC CCGCCCAG CCGCCGCAG ACCGCCATGG AATGTCACGT GTGTGGGTGC CCACATGACC TTTTTGCTGT ACAGCCTTAA TTTTCCTCT ACAGCCTTAA ATCACCTTCT AGTGAACCT AAGAACTGGA ACCACTTGATCC CTTGTACCT TCGGAACCC AAGAACTGGC CCTTGATCC CCTTGATCCC CTTGATCCC CTTGATCCC CCTGACCTCC CCTGACCTCC CCTGACCTCC CCTGCTCCC CCGGCTCC	GGCCCGGCT GGCACCCGGG GGAATACCAG GGATCGTTTTTA GAGGCTACAT GGATCGTCT TGTTAATTCA GGCTGGTAGC AAGAGGATGC TACTCATTCA TCCACGACC GGTGGATCAC GGTCCTTAAA AGAAGGATCA CTGCCCAGCC CTGCCCAGCC CTGCCCAGCC TCAAGTCCCC GGCCTACTT ACAGCTCCT ACAGCTCCT ACAGCTCCT ACAGCTCCT ACAGCTCCT ACAGCTCCT ACAGCTCCT ACAGCTACTT ACCAGTACTT ACCAGTACTT	CCCTGCGCCG AGCAGCCGGG AGCAGCCGAC CCTCTGCAGC CCTCTGGGCC CCTCTGGGCAC CCTGGGCAGC CCTAGGGCAGC CCTAGTGTGT CCAGGTGGAC AGGGTTGATT CAAGGAGGAC CGCCAAGACT CAAAGAGAGAC CCTCATCTGAAAGAGAGC CCTCATGAGAC CCCAAGACT ACAGAAGAGAC CCTCATGACC CCTATAAAGTTG CTACACCGTC CCACATCTCC CCACATCTCC CCACATCTCC CCACATCTCC CCACATCTCC AGCACCCCGC CCACATCTCC CCACATCTCC CCCACACCCGC AGCCCCGCC CCACATCTCC CCACATCTCC CCCACACCCGCC ACCCCCGCC ACCCCCGCC CCACATCTCCC CCACATCTCC CCCACATCTCC CCCACATCTCC CCCACATCTCC CCCACATCTCC CCCACATCTCC CCCACATCTCC CCACATCTCC CCACATCTC CCACATCTC CCACATCTC CCACATCTC CCACATCTC CCACATCT CCACA	CGGCGCGCGC CCCGATCACC CGCCGCCGG GCCGATGGCT TTCACCAAGT TGTTTCCCCT CCTCTCAACA CTCTTCTACT AGCCAACTC AGGAAGGAG GCCTAGCCA CTGTTTCGTG TCCAGAGTCCA GTCCGGGCT ACGTCGGAAC AGGAAGCAG TCCAAGGTGG TCCAAGGTGG TCGAAGGTGG TCGAAGGTGG TCGAAGAGCCCT TTCTTCTTCA CTCATCAAGT TCGTCATCAGTG	51 CGCCGCCGC GCCGCCGG GCCGCCGG GCTTCAGAA TCTACTTCCT AAACCAAAAC CTTTCTGGGA TCTACTCTT TCCTGAGAT TCTACTTCT ACACTCTT ACACTCTT ACACTCTT ACACTCTT ACACTCTT ACGCCAGCC AAGTCGTCC AAGTCGTCT ACGCCAGCC AAGTCGTCC ACGCTGAATGA CTCTGTTTAA AGGCCATCCA TCGTGAATGA TCGTGAATGA TCACTGCCTG GCATGAGGAT TCACTGCCTG GCATGAGGAT	120 180 240 360 420 480 540 660 720 840 900 960 1020 1140 1200 1260 1320 1380
45 50 55 60	CAGGCGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGG	TTGCGGCCCC CCGCCCAG CCGCCCAG ACCGCATGG AATGTCACGT TTTTTGCTGT GCGATATTCC CTTGCTACCT TCTGCTACCT ACAGCCTTAA TTTTCCTCT ACAGCCTTA ATTACCTCT AGTGAACCC AGTGACCCT AGTGACCTCT AAGAACTGGA TCCAAGGAT TCAAGGAT TCCAAGGAT CCATTTTCCG TCCAAGGAT CCAAGGAT CCAAGCTTCG AAGACTGC CCAGACTGC CTCGTGCTGC GTCATTGGGG GTCATTGGGG	GGCCCGGCT GGCAACCGGG CGCTCCGGGG GGAATACCAG GGATCGTTTTA GGAGCTACAT GGATCGTCT TTTTAATTCA GGCTCGTAGC AAGAGGATGC TACTCATTCA TCCACGACCC GGTGCATCAC CTGCCAGACC CTGCCAGACC GGCCTACTT GGCCAGCC TCAAGTCCCC GGCCTACTT GGCCAGACT CCAGTACTT CTGCCAGCT CTGTCTATCG CTGCCAGCT CTGTCTATCC CTGCCAGCT CTGTCTATCC	CCCTGCGCCG AGCAGCCGGG CCCGATCACC CTTCTGCAGC CAACCCGAC CCTCTGGGCAGAC CTGGGCAGAC CTGGGCAGAC CCTGGGCAGAC CCTGGTGTGT CCAGGTGGAC GCTCTTGGC CAAGGTGAC CAAGGAGAC CGCCAAGACT CAAGGAGGAC CCTCATGAGAC CCTCATGAC CCTCATGAC CCTCATGAC CCTCATGAC CCTCATGAC CCTCATGAC CCTCATGAC CCTCATGAC CTACACCCTGC CAAGCCTTG CAAGCCCTTG	CGGCGCGCGC CCCGATCACC CGCCGCCGG GCCGATGGCT TTCACCAAGT TGTTTCCCCT CCTCTCAACA CTCTTCTACT AGCCAACTC AGCAAGCAC CTGTTTCGTG TCCTGTTTCT CCAGAGTCCA AGGAAGCAGC TCCAAGGTGC TCCAAGGTGC TCCAAGGTGC TCCAAGGTGC TGGAACCCT TTCTTCTTCA CTCATCAAGT CTCATCAAGT CTCATCAGTG CTCGTGTTC	51 CGCCGCCGC CGCCGCCGC CGCCGCCGC CCGACCGCCT CCTTCAGAA TCTACTTCCT AAACCAAAAC CTTTCTGGGA TCTAGGTCTTC TCCTGGGATC ACATCACTTT CAGATCGCT ACATCACTT ACGCCAGC AAGTCGTCC AAGTCGTCC CGGTGAAGGT ATGCGAATGA TCTGTTTAA AGGCCATCCA TCGTGAATGA TCACTGCTG CGCTGCATCA TCGTGAATGA TCACTGCTG CGCATGCATTA AGGCCATCCA TCGTGAATGA TCACTGCTG GCATGAATGA TCACTGCTG GCATGAGGAT ATTCAGCCAG	120 180 240 300 360 420 600 660 720 780 840 900 960 1020 1140 1260 1320 1380 1440
45 50 55 60 65	CAGAGGGGGGGGGCGCCGCCGCGCCCGCGCCCCGCGCCCCGCGC	TTGCGGCCCC CCGCGCCAC CCGCCGCGCC ACCGCATGG AATGTCACGT TTTTGCTGT GCATATTCC CTGCTTTCT ACAGCCTTAC ATTTCCTCT TCGGAAACCA ATCACCTTCT AGAGCTTCA AGAACTGAC TCCAAGGATC GCTTTGATCG CTCAAGGATC CCTTGATCGC TCCAAGGATC GCTTTGATCG ATGACCTTTC AAGAACTGGA TCCAAGGATC GCTTTGATCG CTCGTGCTGC CTCGTGCTGC CTCGTGCTGC CTCGTCGGGGG ACGGTCGGGG	GGCCCGGCT GGCTAGCGC AGCAACCGGG GGAATACCAG GGTCCGGGG GGAATACCAG GGTCGTTTTTA GAGGCTACAT GGATCGTCTG TGGCCCAGT TTTTAATTCA TCCACGACCC GGTGGATAC TCCACGACCC GGTGGATCCC TCAAGTCCCT TCAAGTCCC GGCCTACAT AGAGGAATG CTGCCAGC TCAAGTCCC TCAAGTCCC TCAAGTCCC TCAAGTCCC TCAAGTCCC TCAAGTCCC TCACGACTACTT ACCAGTACTT ACCAGTACTT ACCAGTACTT ACCAGTACTT ACCAGTACTT ACGAGTACTT ACCAGTACTT ACGAGTACTT ACGAGTACTT ACCAGTACTT AGGATTGTCAA	CCCTGCGCCG CCCGATCACC CTTCTGCAGC CAACCCCGAC CTTCTGCAGC CTAGATGACA CTGGGCAGAC CTTCTGGTC GCTGGAGAGG CCTAGTGTGT CAAGGTGAGAC CTAGTCTTT TAATCCCTGC AGGGTGAT CAAGAAGAGT CACAGAAGAGT CCTCATGAGC CCTCATGAGC CCTCATGAGC CCTAACACTG CAAGACTG CAACACTGC CAACACTGC CAACACTGC CAACACTGC CAACACTGC CAACACTGC CAACACTGC CAACACTGC CAACACTGC CAACACCTGC CAACACCTGC CAACACCTGC CAACACCTGC CAACACCTGC CAACACCCTG CAACACCCTG CAACACCCTC CAACACCCTC CAACACCCTC CCTCATGTCT	CCGCCGCCGC CCCGATCACC CCCGCCCGG GCCGATGGCT TTCACCAAGT TGTTTCCCT CCTCTCAACA CTCTTCTACT AGGAAGGAG GCCCTAGCCA TCCTTTCGTG TCCGGGGCT ACGTCGGAG TCCAGGTCCA AGGAAGCAGC TCCAAGGTGG TCCAAGGTGG TCCAAGGTGG TCCAAGGTGG TCCAAGGTGG TCCAAGGTGG TCCAAGGTGG TCCAAGGTGG TCCAAGGTGG TCCAAGGTGG TCCAAGGTGG TCCAAGGTGC TCCTCTTCTC TTCTTCTTCA CTCATCACGT GTGATCACCA GTGGACCCTC	51 CGCCGCCGC CGCCGCCGC CGCCGCCGC CCGACCGCCT GCTTTCAGAA TCTACTTCCT AAACCAAAAC CTTTCTGGGA TCTAGGCAT TCAGTCTTC CAGATCACTCTT CAGATCACTCTT ACGCCAGCC AAGTCGTCACTC CGGTTGAAGAT ATCAGATCA TCTGTTTAA AGGCCATCCA TCGTGAATGA TCTGTTTAA TCGTGAATGA TCATGCAGT ATCAGCAGC AAGTCCTGC ATCAGCAGC AAGTCCTGCAATGA TCATGAATGA TCATGAATGA TCATGAATGA TCATGAATGA TCATGAAGAT TCATGAATGA TCATGAAGAT TCATGAAGAT TCATGACCAG AGAGGTTCAT	120 180 240 300 360 420 600 660 720 780 840 900 960 1020 1140 1260 1320 1380 1440
45 50 55 60	CAGGCGGGG GCCGCCGCGC CGCCGCGCC CGCCCGCGCC CTGGGACTGG CACGGTCCTC TGCCTTGGGA AAGAATCGG CACACGCTG AGGGATCATG CAAATTATG CAAAATTATG CTACGTCTAC ACCCTGTTC CCTGTAGGG CCTGGAGGGC TGTTTTTGGTA TGTGACTCAC GGAGGTGGAG GGTGTATAC CACAAGACCC CACAAGACC CAAGACCGCT AAAATCCTC CGACTTGGCC CAAGACCGCT AAAATCCTCC	TTGCGGCCCC CCGCCCAG CCGCCCAG ACCGCCATGG AATGTCACGT GTGTGGGTGC CCACATGACC TTTTTGCTGT GCCATATTCC CTTGCTACCT TCTGCTACCT TCGGAAACCT AAGACCTCT AAGAACTGCA ACGACACT AAGACTGCG CCTGGGTCCC CCAGGATCGC CCAGGATCGC CCAGGATCGC CCAGGATCGC CCAGGTCGGC CCCGTCCTGC ACGTCGGGC ACGTCGGGG ACGTCGAATTA TGGCTGAATC TGGCTGAATC	GGCCCGGCT GGCACCCGGG GGAATACCAG GGATCGTTTTTA GAGGCTACAT GGATCGTCT GGATCGTCT GGCCCAGT TTTTAATTCA GGCTGGTAGC AAGAGGATGC TACTCATTCA TCCACGACC GGTGGATCAC GGCCCAGT CTGCCAGCC GGCCTACTT ACAGTCCC GGCCTACTT ACAGTCCC GGCCTACTT ACCAGTCCT TCAAGTCCC TCAAGTCCT TCGCCACGT ACATGTCTT TCGCCTTC TGGCCCTTC	CCCTGCGCCG CCCGACCCGGC CCAGCCGGC CCAGCCGGC CCTCTGGGCC CTCTGGGCC CTCTGGGCAGC CCTGGGCAGC CCTGGGCAGC CCTGGTGTGT CCAGGTGGAC CCTGGTCTTC CAGGTGGAC CCTCGTCTTC CAGGGTGAC CCCAAGACT CAAAGAGAGA CCTCATGAGC CCTAAAGTG CCACATCTGC CCACATCTCC CCACATCTCC CCACATCTCC CCACATCTCC CCTCATGTCC CCTCCTGCCC CCTCCTGCCC	CGGCGCGCGC CCCGATCACC CGCCGCCGG GCCGATGGCT TTCACCAAGT TGTTTCCCCT CCTCTCAACA CTCTTCTACT AGCCAACTC CTGTTTCGTG GCCCTAGCCA CTGTTTCGTG TCCAGAGTCCA GTCCGGGCT ACGTCGGAAC TCCAAGGTGG TCGAAGCCCT TCTTCTTCA CTCATCAAGT TCGTCATCAGT TTCGTCAGTG GTGATCACCA GTGGACCCCT CTGCAAGTCA GTGGACGCCC CTGCAAGTCA GTGGACGCCC CTGCAAGTCA GGAGTGCCGC	51 CGCCGCCGC CGCCGCCGC CGCCCCGC CCGACCCGC CCGACCCGC CCGACCCGC CCGACCCGC CCTTCAGAA TCTACTTCT AAACCAAAAC CTTTCTGGGA TCTTCTGGGA TCTACATCTT CAGATCGCTT CAGATCGCTT ACCGCCAGCC AAGTCGTCTT ACCGCCAGCC AAGTCGTTCTAA AGGCCATCCA TCGTGAATGA CTCTGTTTAA AGGCCATCCA TCGTGAATGA TCACTGCCTG GCATGAGGAT ATCACCCGG GCATGAGGAT ATTCACCCAG AGAGGTTCAT TCCTTGCTCT TCGATGGTCCT	120 180 240 300 360 420 540 600 720 780 840 960 1020 1140 1200 1140 1320 1380 1440 1500 1560 1620
45 50 55 60 65	CAGGCGGCG GCGCGCGCGCGCGCGCGCGCGCGCGCGCGC	TTGCGGCCC CCGCCCAG CCGCCCAG CCGCCCAG ACCGCATGG AATGTCACGT CTTTTGCTGT GCGATATTCC CTTGCTACCT CTCACTTTCT ACAGCCTTAA TTTTCCTCT ACAGCCTTA ATTACCTCT AGTGAACCC AGGACTGC AGGACTGC CCTGGCTGC CTCGTTGGTGC CCAGACTGC CTCGTGCTGC CTCATTGGGG ACGTCGGGGGA ACGTCGGGGGGA CGTACATTC CGCAGACTCC TCGTGCTGC CTCATTGGGG ACGTCGGGGG ACGTCGATCC CTCAATGCTC	GGCCCGGCT GGCAACCGGG GGAATACCAG GGATCGTTTTTA GAGGCTACAT GGATCGTCT TTTTAATTCA GGCTGGTAGC AAGAGGATGC TACTCATTCA TCCACGACCC GGTGGATCACC GGTCCTTAAA AGAAGGATGC TCAAGACCC GGCCTACTT GGCCCAGCC TCAAGTCCCC GGCCTACTT ACGACTACTT ACCAGTACTT ACCAGTACTT ACCAGTACTT TCTCTATCG AGATTGTCAA ACATGATCTT TGGCCCATT	CCCTGCGCCG AGCAGCCGGG CCCGATCACC CTTCTGCAGC CAACCCGAC CCTCTGGGCAGC CTTGGGCAGAC CTTGGGCAGAC CTTGGGCAGAC CTTGGGTGTGT CCAGGTGGAC GCTAGTGTGT CAAGGAGGAC CGCCAAGACT GAAGAGGAG CCTCATGAGAG CCTCATGAGC CTAAAGTGTG CTAAACTTG CAAGCCCTG CAAGCCTG CCAAGCTT CTAAAGTTG CCACATCTGC CCACATCTGC CGAAGCCTG CCTCATGACC CTCATGACCCTG CGTCCTGGCT GAAGACCAAG	CGGCGCGCGC CCCGATCACC CGCCGCCGG GCCGATGGCT TTCACCAAGT TGTTTCCCCT CCTCTCAACA CTCTTCTACT AGCAAGTCA AGGAAGGAG GCCTAGCCA CTGTTTCTG TCCTGTTTCT CCAGAGTCCA AGGAAGCAGC TCCAAGGTGC TCCAAGGTGC TCCAAGGTGG TGGAACCCT TTCTTCTTCA CTCATCAAGT CTGCTGTTTC CTGCTGTTTC CTGCTGTTTC CTGCTGTTTC CTGCTGTTTC GTGATCACCA GTGGACGCC CTGCAAGTCA GTGGACCCA GTGGACGCCA CTGCAGCTCA	51 CGCCGCCGC CGCCGCCGC CGCCGCCGC CCGACCGCC CCGACCGCCT GCTTTCAGAA TCTACTTCT AAACCAAAAC CTTTCTGGGA TCTAGGCAT TCAGACTCT CAGATCTCT ACATCACTT CAGATCGCC ACATCACTT CAGATCGCC CGGTGAAGGT ATGCGAATGA TCATGGCAATGA TCACTGCTTAA AGGCCATCCA TCGTGAATGA TCACTGCTG GCATGAATGA TCACTGCTG ATTCAGCAG AGAGGTTCAT TCCTTGCTCT TGATGGTCCT TGGCCCACAT	120 180 240 300 360 420 660 660 720 780 840 900 1020 1140 1200 1260 1320 1440 1560 1560 1620 1680
45 50 55 60 65	CAGGCGGGG GCCGCCGCGC CGCCGCGCC CGCCCGCGCC CTGCACTGC CAGGTCCTC CTATCTCTCC TGCCTTGGGA AAGAAGTCGG CACACGCTG AGGATCATG CACACGCTG ACCCTGTTC CCTGCAGGG CCTGCAGGG CTGTTACTC GGAGGTGGAG GGTGTTATAC CGACCTGATG CACGAAGGCC CAGAACGCT AAAATCCTCC GGACTTGCGC CAGACCGCT AAAATCCTCC GGACTTGCC CAACACCGCT AAAATCCTCC GGACTTCGCC CTACCTCCTG GACTTCCCTC GACACACAAA	TTGCGGCCC CCGCGCCC CCGCGCCC CCGCGCCC ACCGCATGG AATGTCACGT TTTTGCTGT GCATATTCC CTGCTTTCT ACAGCCTTAC ATTTCCCTCT TCGGAAACCA ATCACCTTCT AAGAACTGA TCCAAGGATC GCTTTGATCG CTGTGACCTCT CTGGGAACCA ATCACCTTTT CGGAAACTGA TCCAAGGATC CCAGACTGC CTCATTTCCGGCTGC CCAGACTGC GTCATTGGGG ACGTCGGGG ACGTACATTA TGGCTGATCG GACAATCGGA CGTCAATCGGGG ACGTACATTA TGGCTGATCTG GACAATCGGA	GGCCCGGGCT GGCTAGCGGG GGAATACCAG GGCTCCGGGG GGAATACCAG GGATCGTTTTTA GAGGCTACAT GGATCGTTCA TTTTAATTCA GGCTGGTAGC AAGAGGATGC TACTCATTCA TCCACGACC GGTGGATCAC GGTGGATCAC TCAAGTCCC GGCCTACAT ACGACCC GGCCTACAT ACGACCC GGCCTACAT ACGACCC TCAAGTCCC TCAAGTCCC TCAAGTCCC TCAAGTCCC TCAAGTCCC TCAAGTCCC TCAAGTCCC TCAAGTCCC TCAAGTCCT TCAAGTCCT TCAAGTCAT TCAAGCCGAC TCAAGCCTGAT TCAAGCCGAT TCAAGCCGAT TCAAGCCGAT TCAAGCCGAT	CCCTGCGCCG CCCGATCACC CTTCTGCAGC CAACCCCGAC CTTCTGCAGC CTCTGGGCC TCAGATGACA CTGGGCAGAC CTTTCTGGTC GCTGGAGAGG CCTAGTGTGT CAAGGTGAC CTAGTCTTT CAAGGAGAGA CTCATCACC CAAGACAC CACAAGACT CAAGAAGAGAC CTTAAACTTG CTACACCGTG CAAGACCTG CAAGACCTG CAAGACCTG CAAGACCTG CAAGACCTG CAAGACCTG CAAGACCTG CAAGACCTG CAAGACCTG CAAGACCAG GAAGACCAG GAACCAAGACT GAAGACCAG GAACCAAGACT	CCGCCGCCGC CCCGATCACC CCCGCCCGG GCCGATGGCT TTCACCAAGT TCTACCAAGT CCTCTCAACA CTCTTCTACT AGGAAGGAG GCCTAGCCA CTGTTTCGTG TCCAGAGTCCA AGGAAGCAC TCCAGCAGC TCCAGCAGC TCCAGCAGC TCCAGCAGC TCCAGCAGC TCCAGCGGAC TCCAAGGTGG TGGAACCCT TTCTTCTTC CTCATCAGT CTGCTACTCA GTGGTGTTTG GTGGTGTTTG GTGGTGTTTG GTGGTG	51 GGCGCCCCC CGCCGCCCGC CGCCCCCGC CCGCCCCCC	120 180 240 300 360 420 540 600 660 720 780 840 900 1020 1140 1260 1320 1320 1340 1560 1560 1680 1740
45 50 55 60 65 70	CAGGCGGGG GCCGCCGCGC GCCGCGCGC CGCCCGCGCC CTGCACTGG CACGGTCCTC TGCCTTGGA AAGAATCATG CAAAATTATG CAAAATTATG CAAAATTATG CTGCTTGGAG CCTGCAGGG CTGCAGGGC TGTTTTGGTA GGAGTGATAC CGACCTGATAC CGACCTGAAGCC CACAAGACCGCT AAAATCCTC CAGAAGCCC CAAGACCGCT AAAATCCTC CGACTTTAGAC CACTGATAC CGACTTGATAC CACCAAGACCC CAAGACCGCT AAAATCCTCC CAAGACCGCT AAAATCCTCC CGACTTGCCC CTACCTCCTG CATGGTGCC CATGCTACAAAAAACCTTTAT	TTGCGGCCCC CCGCCGCAG CCGCCGCGCAG ACCGCCATGG AATGTCACGT GTGTGGGTGC CGACATGACC TTTTTGCTGT GGCATATTCC CTCACTTTCT ACAGCCTTAA ATTTCCCTCT AGGAACCA ATCACCTTCT AAGAACTGGA ATCACCTTCT AAGAACTGGA CCTTGATCG CCTGGTCTGC CTCATTTCCC CCAGACTGCC CTCATTTCCC CCAGACTGCC CTCATTGGGG ACGTCGCGC ACGTCGCGC ACGTACATTA TGCCTGAATC GCTAATGCGA CGCAAATCGGA CCCTGGGAGC CCCTGGGAGC CCCTGGGAGC CCCTGGGAGC CCCTGGGAGC CCCTGGGAGC CCCTGGGAGC CCCTGGGAGC CCCTGGGAGC	GGCCCGGCT GGCTAGCGC AGCAACCGGG GGAATACCAG GGCTCCGGGG GGAATACCAG CTTGTTTTTA GAGGCTACAT GGATCGTCT GGCCCACT AGAGGATGC TACTCATTCA TCCACGACCC GGTGGATCAC GGTGGATCAC GGTGGATCAC GGTGGATCAC GGTGGATCAC GGTGGATCAC GGTCCTTAAA AGAGGATGC TCACGACCC TCAGTCCC GGCCCTACTT ACCAGTACTT ACCAGTACTT ACCAGTACTT TCTGTCTATCG TGGCCCTTC TGATGGCGAT TCAGGCCCTTC TGATGGCGAT TCAAGCTGAT TCAAGCTGAT TCAGCTGAT TCAGCTGAT TCAGCTGAT TCAGCTGAT TCAGCTGAT TCGCATTCAA	CCCTGCGCCG AGCAGCCGGG CCCGATCACC CTTCTGCAGC CAACCCCGAC CTCTGGGCC TCAGATGACA CTGGGCAGAC CTGGGCAGAC CTAGTGTGT CCAGGTGGAC CCTAGTGTGT CAAGGAGAC CCTAGTGTGT CAAGGAGAC CCTCATCTGC AGGGTTGATT CAAGGAGGAC CCTCATGAGC CCTCATGAGC CCTCATGAGC CCTCATGAGC CCTCATGAGC CCTCATGAGC CCTCATGAGC CCTCATGTC CCACACCTGC CAAGGCCCC CGTCCTGGCT CGAAGACCAAGACT GAAAGACAAAGTT GGACAAGGTT GGACAAGGTT	CGGCGCGCGCGCGCCCGGTCACCCGCGCGCGGGGCTCACCAAGTCACAAGTCGGAGGCGCTCAAGCAAG	51 GGCGCCCCC CGCCGCCCGC CGCCCCCGC CCGCCCCCC	120 180 240 300 360 420 660 660 720 780 840 900 1020 1140 1200 1260 1320 1440 1560 1560 1620 1680
45 50 55 60 65	CAGGCGGGG GCCGCCGCGC CGCCGCGCC CGCCGCGCC CGCCGC	TTGCGGCCCC CCGCGCCCC CCGCGCCCC CCCCCGCGCC ACCGCCATGG AATGTCACCT TTTTGCTGT GCATATTCC CTGCTACTTCT ACAGCTTAC ATTCCCTACT TCGGAAACCA ATCACCTTCT AAGAACTCA ATCACCTTTT AAGAACTCGA ACCAAGGATC CTGGGAACCA ATCACTTTT AGTACTTTT AGTACTTTT AGTACTTTT AGTACTTTT AGTACTTTT CGAAGCTTTG AAGACTTTG AAGACTTTG CTGACTGGC CTCAGCTGGC ACGTCGGG ACGTCGGG ACGTCGGG ACGTCGGG ACGTCAATGCT GTCAATGCT GACAATCCGA GCCTGGGAGC CTGAAGAACT CTGATGGCC	GGCCCGGGT GGCAACCGGG GGAATACCAG GGCTCCGGGG GGAATACCAG GGATCGTTTTTA GAGGCTACAT GGATCGTTCT TTTTAATTCA GGCTGGTAGC AAGAGGATGC TACTCATTCA TCCACGACCC GGTGGATCACC GGTGGATCACC GGCCTACTT ACAGTACT ACAGTACTT ACCAGTACTT ACCAGTACTT ACCAGTACTT ACCAGTACTT TCGATGCCTTCA ACATGATCT TGGGCCTTCT TGATGGCGAT TCAGCCTTCAT TGATGGCGAT TCAGCCTACT TGATGCACATT TGTGCACATT TGTGCACATT	CCCTGCGCCG CCCGATCACC CTTCTGCAGC CAACCCCGAC CTTCTGCAGC CTAGATGACA CTGGGCAGAC CTGGGCAGAC CTGGGCAGAC CTGGGCAGAC CTGGGCAGAC CTGGGCAGAC CTAGTGTCT CAAGGTGAC CAGGTGAC CAGAGACAC CAGAAGACT CAAGAAGAC CTCATCAGC CTCATCAGC CTCATCTCT CAAGACCTTC CAACACTTCC CACACTCTC CCACACTCTC CCTCATCTCT CTCAGCCCCC CGTCTCTCGCC CGAGACAATT GGAACAAGTT GGAACAAGTT GGAACAAGTT GGAACAAGTT GGAACAAGTT GGAACAAGTT CGACCGTC CTCAGCCCTC CTCACCCTC CTCACCTC CTCACCCTC CTCACCC CTCACC CT	CCGCCGCCGC CCCGATCACC CCCGCCCGG GCCGATGGCT TTCACCAAGT TGTTTCCCCT TGTTTCCCT AGCAAGTCA AGGAAGGAG GCCTAGCCA TCCTGTTTCT CCAGAGTCCA GTCTGTTTCT CCAGAGTCCA AGGAAGCAC TCCTGTTTCT TCCTCGGGGCT TTCTTCTTCA TCCAGAGTCCA TCCAAGTTG TTCTTCTTCA TTCTTCTTCA TTCTTCTCA TTCTCTCATC TTCTCTCAGT TTCTCTCAGT TTCTCTCAGT CTGCAAGTC GTGACCCT CTGCAAGTC GTGACCCC GTGACGCG CTGCAAGTCA GGAGTGCCG CTGCAAGTCA GGAGTGCCG CTGCAATCA GGACTCCC CTGCAATCA GGACTCCC CTGCAATCA GGACTCCC CTGCAATCA GGACTCCA CTGCCATCA GCACCTTCA GTGACCATTC	51 CGCCGCCGC CGCCGCCGC CGCCGCCGC CCGCCCGC	120 180 240 300 360 420 660 660 720 780 840 900 1020 1080 1200 1260 1380 1440 1500 1680 1740 1860 1860 1920
45 50 55 60 65 70	CAGGCGGGG GCCGCCGCGC CGCCGCGCC CGCCCGCGCC CTGCACTCC CTATCTCTCC TGCCTTGGGA AAGAAGTCGG CACCACGCTG AGGATCATG CAAAATTATG CTACGTCTAC CTGCTGGAGGC CTGCTGGAGGC CTGCTGGAGGC CTGCTGATG CGACCTGATG CACCAGACCGC CAAGACCGC CAAGACCGC CAAGACCGC CAAGACCGC CAAGACCGC CAAGACCGC CAAGACCGC CAAGACCGC CAAGACCAAA AAAGCTTTAT GCTGAAGGTG CATGGTGCC CAAGACCAAA AAAGCTTTAT CCTGCAAGGTG CATGGTGCC CAAGACCAAA AAAGCTTTAT CCTGAAGGTG CACGCCCTTT	TTGCGGCCCC CCGCGCCCC CCGCCGCCCC CCGCCGCCCC ACCGCGCCC ACCGCCATGG AATGTCACGT TTTTGCTGT GCCATATTAC CTTGCTACCT CTCACTTTCT ACAGCCTTAA ATTTCCCTCT TCGGAAACCA ATCACCTCT AGAACTGA TCCAAGGATC CCAGACTGC CTCGTCTCC CTCACTTCT AGAACTGA ACCATTCT AGAACTGA ACGACTGC CTCGTCCTC CTCATCGGG ACGTCGCGG ACGTACATTA TGCCTGAATCC GTCAATCGGA CCTGGAATC GTCAATCGGA CCTGGAACCT CTCAATCGGA CCTGGGAGC CTCAATCGGA CCTGGAACCGC CCCAGACACA	GGCCCGGCT GGCTAGCGC AGCAACCGGG GGAATACCAG GGATCGTATTTA GAGGCTACAT GGATCGTCT TGGCCCACGT TATTAATTCA GGCTGGTAGC AAGAGGATGC TACTCATTCA TCCACGACCC GGTGGATCAC GGTGGATCAC TCAGGACCC TCAAGTCCC TCAAGTCCT TCTCTATCG TCAAGCCGTTCAA ACATGATCTC TCATGCGCAT TCAAGCTGAT TCAAGCTGAT TCACCTACCT TCTGCCACATT CCTTCCTTCTTCTTCT TCTCTCACTTCC TCTCCTACCT	CCCTGCGCCG AGCAGCCGGG CCCGATCACC CTTCTGCAGC CAACCCCGAC CTCTGGGCC TCAGATGACA CTGGGCAGAC CTTTCTGGTC GCTGGTGTGT CCAGGTGGAC GCTCGTCTTG TAATCCCTGC AGGGTTGATT CAAGGAGGAC CGCCAAGACT GAAAGAGAGT CCTATAGTC CTACACCGTC GAAGGCCCT GAAGGCCCT GAAGGCCCT GAAGGCCCC GTCCTGGCT GAAGACAAGTT GCACAAGAT GGACAAGAT TCAGCCCC CTCATGACC CTCATGACC GTCAGCCCC CTCATGACC GTCAGCCCC CTCAGGCCTC CTCAGCCCC CGTCCTGGCT GAAGACAAATT GGACAAGGTG TCAGCCCTTA	CCGCCGCCGC CCCGATCACC CCCGCCCGG GCCGATCGCT TTCACCAAGT TTCACCAAGT CCTCTCAACA CTCTTCTACT AGCAAGGAG GCCTAGCCA CTGTTTCGTG TCCAGGAGCCA TCCAGGAGCCA TCCAGGAGCCA TCCAGGAGC TCCAGGAGC TCCAGGAGC TCCAGGAGC TCCAAGGTGG TGGAACCCT TTCTTCTTCG TTGGTCAGTG GTGATCACA GTGGACCCT CTGCAGTCAGCA CTGCTGTTTCG CTGCTGTTTCG CTGCTGTTTCG CTGCTGTTTCG CTGCTGTTTCG CTGCTGTTTCG CTGCTGTTTCG CTGCTGTTTCG CTGCTGTTTCG CTGCAGTCA GGAACCTCA GGAACTCAGG CTCAATGGGA CTCAATGGGA CTCACATCA GGCACCTTCA GGCACCTTCA TTCAACATCC	51 GGCGCCGCC CGCCGCCGC CGCCGCCGC CGCCGCCGC CCGACCCGC CCGACCCGC CCGACCCGC CCGACCCGC CCGACCCGC CCGACCCGC CCGACCCGC CCGACCCGC CCGACCAAAAC CTTTCTGGGA TCTTCTGGGA TCTTGGCAT TCAGTCTT ACACACTT ACACACTT ACACACTT ACCGCCAGC AAGTCGTTCCT ACGCCAAGC CAGGGAATGA TCACTGCTG GCATGAAGGA TCACTGCTG GCATGAGGAT ATTCAGCCAG AGAGGTTCAT TCCTTGCTCT TGATGGTCCT TGATGGTCCT TGAAGGACAA TCCGGGTCTG GCAGGAGGA CCTGGGTCTG GCAGGAGGA TCCGGGTTCTC TCAAAGTCCT GCACGGACCA TCCAGGACCAA TCCGGGTTCTC	120 180 240 300 360 420 540 600 660 720 780 840 900 1020 11200 1260 1320 1320 1350 1440 1500 1560 1680 1740 1800 1800 1920 1980
45 50 55 60 65 70	CAGGCGGGG GCGCCGCGCGC GCGCCGCGCC CGGCCGCGCC CGGCCGCGCC CTGGGACTGG CAGGTCTTC TGCTTGGA AAGAATCAG CAAAATTATG CAAAATTATG CAAAATTATG CCTGCAGGG CCTGCAGGG CCTGCAGGG GGTGTTATAC CGACCTGATA CGACCTGATA CAACACCGCT AAAATCCC CAAGACCGC CAAGACCGC CAAGACCGCCT AAAATCTCC CGACTGCAGC CAAGACCGCCTTA CAACACCTCTT CATCTCTCC CAAGACCCCATTC CAACACCCCTTTT CATCCTCGAAGGCC CAAGACCCCCTTT CAACACCCCTTTT CATCCTGGATC CACGCCCTTTT CATCCTGGATC CACGCCCTTTT CATCCTGGATC CACGCCCTTTT CATCCTGGATC CACGCCCTTTT CATCCTGGATC CACGCCCTTT CATCCTGGATC CACGCCCTTT CATCCTGGATC CATCCTGGATC CATCCTGGATC CATCCTGGATC CATCCTGGATC CATCCTGGATC CATCCTGGATC CATCCTGGATC CCTGAACATT	TTGCGGCCCC CCGCCGCAG CCGCCGCAG ACCGCCATGG AATGTCACGC TTTTTGCTGT GGCATATTCC CTTCATTTCT ACAGCTTAT ATTTCCCTCT TCGGAAACCA ATCACCTTCT AAGAACTGGA ATCACCTTCT AAGAACTGA ATCTTTTCATTC CCAGAACTGC CTTGTTTTCATCG CAGACTGC CTCATTTTCC CAGACTTCT AAGACTGGC CTCATTGGGG ACGTCGGGGACGC CTCATGGGGG ACGTACATTA TGGCTGAATC GCCAGAATCGA CCTCAATGCTG CTCAATGCTG GACAATCGGA CCTCAGGAGC CTCAGGAGC CTGGGGGC CTGAAGAACT CTGGTGGCCT CTGGTGGCCT CTGGTGGCCT CTGCTGGCACAC CTCCCAGACAG CTCCCATGG	GGCCCGGCT GGCTAGCGC AGCAACCGGG GGAATACCAG GGATCCTGGGG GGATCCACT GGACCCACT TTTTAATTCA GGCTGGTAGC AAGAGGATGC TACTCATTCA TCCACGACCC GGTGGATCAC GGTCCTTAAA ACAGACCC GGCCCACT TCAGCCC GGCCTACTT ACCAGCCC TCAGTCCT TCAGCTCT TCAGCTCT TCAGCTCT TCAGCGCT TCAGCGCT TCAGCCCT TCAGCCT TCATCAGCAC TCAGCCT TCATCAGCAC TCAGCCTACCT TCAGCCTACCT TCAGCCACC TCATCAGCAC TCATCAGCAC	CCCTGCGCCG AGCAGCCGGG CCCGATCACC CTTCTGCAGC CAACCCCGAC CTTCTGGTC CTCGGGCAGC CTTGGGCAGC CTTGGGCAGC CTTGGGCAGC CTTGGGCAGC CTTGGTGTGT CAGGTGGAC CCTCGTCTTG AGGGTGAT CAAGGAGGAC CGCCAAGACT CGAAGAGGAC CTTAAAGTTG CTAACCGTG CACATCTGC GAAGGCCCC CGCCATGTC CTCATGTGC CACATCTGC CACATCTCAC CACCCTCTACC CTCAGCCCC CGTCCTGCCT CACCCTTCACC CTTCAGCCCTC CATCGTCTAC CATCTGCCTTGC CATCGTCCAC CATCGTGCACC CATCTGCCACC CATCGTGCACC CATCTGCACC CATCGTGCACC CATCTGCACC CATCGTGCACC CATCTACC CATCGTGCACC CATCGTGCACC CATCGTGCACC CATCGTGCACC CATCGTGCACC CATCGTGCACC CATCGTGCACC CATCGTGCACC CATCGTGCACC CATCATCTACC CATCGTGCACC CATCATCTACC CATCGTGCACC CATCATCACC CATCATCACC CATCATCTACC CATCATCACC C	CGGCGCGCGC CCGATCACC CCGCTCGCGG GCCGATGGCT TTCACCAAGT TTCACCAAGT TCTTCTACT AGCCCAACT AGCCCAACT CCTTTCTACT AGCAGGAG GCCTAGCCA CTGTTTCGTG TCCAGGAGTCCA GTCGGAGCCA CTGTTTCTTC TCAGGAGCCT ACGTCGGACC TTCTTCTTCT TCATCAAGT TCGTCATTTC TTCATCAAGT TTCGTCAGTG TTCGTCACTG GTGACCCTT GTGACCATC GAGTGCGA CCTGCAAGTCA GGAGTGCG CTGCAAGTCA GGAGTGCG CTGCAAGTCA GGAGTGCCATCA GGACCTTCA GCGCCATCA GCGCCTTCA GCGACCTTCA GCGACGTGTT TTCACCATC CTGCACATCC CCGAGTGCCT TTCACCATCA GCGACCTTCA GCGACGTCT CCCGAGTGCCT TTCACCATCC CCGAGTGCCT TTCACCATCC CCGAGTGTCT	51 CGCCGCCCCC CGCCGCCCCC CGCCGCCCGC CCGACCCGC CCGACCCGC CCGACCCGC CCGACCCGC CCGACCCGC CCGACCCGC CCGACCCGC CCGACCCGC CCGACCCCA AACCAAAAC CTTTCTGGGA TCTTCTGGGA TCTTCTGGCAT TCAGTCTTC ACATCACTTT ACCGCCAGCC AAGTCGTCCT ACCGCCAGCC AAGTCGTCATTAA AGGCCATCCA TCATGATTAA AGGCCATCCA TCATGATTAA TCATGCCTG GCATGAATGA TCACTGCCTG GCATGAGTCAT TCATGCTCT TGATGGTCCT TGATGGTCCT TGATGGTCCT TGATGGTCCT CGCAGAGAA CCTGGGTTCC CCCTCAAACG CCCCCCCAAACG CCCCCCCAAACG CCCCCCCAAACG CCCCCCCC	120 180 240 300 360 420 540 660 720 780 840 900 1020 1080 1140 1200 1320 1380 1440 1500 1560 1620
45 50 55 60 65 70	CAGGCGGCG GCGCGCGCGCGCGCGCGCGCGCGCGCGCGC	TTGCGGCCCC CCGCCCAG CCGCCCAG ACCGCCATGG AATGTCACGT GTGTGGGTGC CCACATGACC TTTTTGCTGT GGCATATTCC CTTGCTACCT TCGGAACCAT ACACCTTCT ACAGACTCT AGGAACCGC ATGTTTTCCTCT AGGAACTGGA TCCAAGGAT CCAAGGAT CCAAGGAT CCAGACTGG CCTGGTGCTGC GCAGACTGGC GTCATTGATCG GACATCGGC CTGGTGCTGC GCCAGACTGG CCCAGACTGG CCCAGACTGG CCCAGACTGC CCCAGACTGC CCCAGACTGC CCCAGACTGC CCCAGACTGC CCCAGACTGC CCCAGACTGC CCCCAGACTGC CCCCAGACTGC CTCCCAGCTGC TTTCTCCCC	GGCCCGGCT GGCACCGGG GGAATACCAG GGATCGTTTTTA GAGGCTACAT GGATCGTCT GGATCGTCT GGATCGTCT GGCCCAGT TTTTAATTCA GGCTGGTAGC AAGAGGATGC TACTCATTCA TCACGACCC GGTGGATCAC GGCCCAGT CTGCCCAGCC TCAGTCCC GGCCTACTT ACCAGTCCC GGCCTACTT ACCAGTACTT ACCAGTACTT ACCAGTACTT TCAGTACTT TCAGTACTC TGGCCCTTC TGATGGCGAT TCAAGCTCCC TGATGGCGAT TCAAGCTCCC TGATGGCGAT TCAAGCTCCC TGATGGCGAT TCAACCTTC TGGCACTTC TGGCCTTCCTTCCTTC TGTCCACCT TCATCACCAC ATGAGGAGCT TCATCAGCAG ATGAGGAGCT TCATCAGCAG ATGAGGAGCT TCATCAGCAG ATGAGGAGCT TCATCAGCAG ATGAGGAGCT TCATCAGCAG ATGAGGAGCT TCATCAGCAG ATGAGGAGCT ACGCATCACAC ATGAGGAGCT TCATCAGCAG ATGAGGAGCT ATGAGGAGCT ACGCATCAC ATGAGGAGCT ACTCACCACAC ATGAGGAGCT ACGCATCAC ATGAGGAGCT ATGAGGAGCT ACGCATCAC ATGAGAGAGCT ACGCATCAC ATGAGGAGCT ACGCATCAC ATGAGGAGCT ACGCATCAC ATGAGAGAGCT ATGAGGAGCT ACGCATCAC ATGAGGAGCT ACGCATCAC ATGAGAGAGCT ATGAGAGAGCT ACGCATCAC ATGAGAGAGCT ATGAGGAGCT ATGAGAGAGCT ATGAGAGAGACT ATGAGAGAGCT ATGAGAGAGACT ATGAGAGAGCT ATGAGAGAGACT ATGAGAGAGCT ATGAGAGAGACT ATGAGAGAGAACT ATGAGAGAGAGACT ATGAGAGAGACT ATGAGAGAGACT ATGAGAGAGAGACT ATG	CCCTGCGCCG AGCAGCCGGG CCCGATCACC CTTCTGCAGC CAACCCGAC CTTCTGGGC TCAGATGACA CTGGGCAGAC CTTGGGCAGC CCTAGTGTGT CCAGGTGGAC CCTGGTCTTG CAGGTGGAC CCTCGTCTTG CAAGGAGGAC CTATCTCTGT CAAGGAGGAC CTCATCTTC CAAGGAGGAC CTCATGAC CTCATGAC CTCATGAC CTCATCTGC CAACTCTG CCACATCTGC CCACATCTGC CCACATCTGC CCACATCTGC CCACATCTGC CCACATCTGC CTCATGACC CTCATGACC CTCATGCCCT GAAGACCAAG CTCAAGCCTT GCACACCTGC CTCACTCTC CACACCTGC CTCACTCTC CACACCTGC CTCACCCTC CACACCTCC CCTCCTGCCT CACACCTCT CACACCTTC CACACCTTC CACACCTTC CACACCTTC CACACCTTC CACACCTTC CACACCTTC CACACCTTC CACCCTTC CACCTTC CACCCTTC CACCTTC CACCCTTC CACCCTTC CACCCTTC CACCCTTC CACCCTTC CACCCTTC CACCTTC CACCCTTC CACCCTTC CACCCTTC CACCCTTC CACCCTTC CACCTTC CACCCTTC CACCCTTC CACCTTC CACCCTTC CACCTTC CACCCTTC CACCTTC CAC	CGGCGCGCGC CCGATCACC CGCGGCCGGG GCCGATGGCT TTCACCAAGT TTCACCAAGT CTCTTCTACT AGCCCAACTC CTCTTCTACT AGCCCAACTC CTGTTTCGTG TCCAGGAGGAG GCCTAGCCA CTGTTTCGTG TCCAGAGTCCA GTCAGGAGCCA TCTTCTTCTTCT AGCTCGGACC TTCTTCTTCA TCGAGAGTCCA TCGACACTC TTCTTCTTCA TCGCACACT TTCTCTCACACT GTGACCCT CTGCAAGTCA GGAGTGCG GCACTTCA GCGCACTTCA GGGACCTTCA GCGCACTTCA GCGCACTTCA GCGACTTCC GCGGGGTGTCT TTCACACACC GCGAGTGTCT TTCACACACC GCGACTTCA GCGACTTCA GCGACTTCA GCGACTTCA GCGAGTGTCT AGCATCACACC GCGAGTGTCT AGCATCACACC GCGAGTGTCT AGCATCACACC GCGAGTGTCT AGCATCACACC GCGAGTGTCT AGCATCACC	51 GGCGCCGCC CGCCGCCGC CGCCGCCGC CGCCGCCGC CCGACCCGC CCGACCCGC CCGACCCGC CCGACCCGC CCGACCCGC CCGACCCGC CCGACCCGC CCGACCCGC CCGACCAAAAC CTTTCTGGGA TCTTCTGGGA TCTTGGCAT TCAGTCTT ACACACTT ACACACTT ACACACTT ACCGCCAGC AAGTCGTTCCT ACGCCAAGC CAGGGAATGA TCACTGCTG GCATGAAGGA TCACTGCTG GCATGAGGAT ATTCAGCCAG AGAGGTTCAT TCCTTGCTCT TGATGGTCCT TGATGGTCCT TGAAGGACAA TCCGGGTCTG GCAGGAGGA CCTGGGTCTG GCAGGAGGA TCCGGGTTCTC TCAAAGTCCT GCACGGACCA TCCAGGACCAA TCCGGGTTCTC	120 180 240 300 360 420 600 660 720 780 840 900 960 1020 1140 1260 1380 1440 1500 1560 1680 1740 1860 1920 1980 2040 2100

	CGTGGTGGGC	CAGGTGGGCT	GCGGAAAGTC	GTCCCTGCTC	TCAGCCCTCT	TGGCTGAGAT	2280
		GAGGGGCACG					2340
						TGGAGGAACC	2400
5						TCCTGCCCAG	2460
J						AGAAGCAGCG	2520
						ATGATCCCCT	2580
						GCCCCAAGGG	2640
						TGCCGCAGGT ACCAGGAGCT	2700
10						CAGAGCAGGA	2760 2820
10						AAGCAAAGCA	2880
						GACAGCTCAG	2940
						CAGAACTGCA	3000
						AGGCGCAGAC	3060
15	AGGGCAGGTC	AAGCTTTCCG	TGTACTGGGA	CTACATGAAG	GCCATCGGAC	TCTTCATCTC	3120
	CTTCCTCAGC	ATCTTCCTTT	TCATGTGTAA	CCATGTGTCC	GCGCTGGCTT	CCAACTATTG	3180
						CGAAAGTCCG	3240
						GCTACTCCAT	3300
20						TGCTGCACAG	3360
20						TGGTGAACCG	3420
						AGATGTTCAT	3480
						CGCCCATCGC	3540 3600
						ATTCCCATTT	3660
25						AGCGCTTCAT	3720
						GCATCGTGGC	3780
		CTGGCCGTGC					3840
	CCTGTTTGCG	GTGATCTCCA	GGCACAGCCT	CAGTGCTGGC	TTGGTGGGCC	TCTCAGTGTC	3900
20	TTACTCATTG	CAGGTCACCA	CGTACTTGAA	CTGGCTGGTT	CGGATGTCAT	CTGAAATGGA	3960
30						AGGAGGCGCC	4020
						GAGTGGAATT	4080
		TGCCTGCGCT					4140
		GGGGGAGAAA					4200
35						TCATCGATGG TCATCCCCCA	4260 4320
33		TTGTTTTCGG					4380
						TGTCAGCCCT	4440
		CTAGACCATG					4500
40						TGTTGGATGA	4560
40						TCCGGACACA	4620
						TGGACTACAC	4680
						CGGACCTCCT	4740
						GAGCCCCAGA	4800
45						GGAGTCAGTA	4860
73						CCCAGACAAC CTGGCTGTGA	4920 4980
		GAGACAGAGA			CIGCCIGGAA	CIGOCIGIGA	4360
				•			
50		BH2 Protein seque	ence:				
50	SEQ ID NO:22 O	8H2 Protein sequents: AAB46					•
50	SEQ ID NO:22 O Protein Accession	#: AAB46	616	31	41	51	
	SEQ ID NO:22 O		616 21	31 	41 	51 !	
50 55	SEQ ID NO:22 Q Protein Accession	1#: AAB46	616 21 	1		51 PFYFLYLSRH	60
	SEQ ID NO:22 O Protein Accession 1 MALRGFCSAD DRGYIOMTPL	#: AAB46 11 GSDPLWDWNV NKTKTALGFL	21 TWNTSNPDFT LWIVCWADLF	 KCFQNTVLVW YSFWERSRGI	 VPCFYLWACF FLAPVFLVSP	 PFYFLYLSRH TLLGITTLLA	60 120
	SEQ ID NO:22 Q Protein Accession MALRGFCSAD DRGYIQMTPL TPLIQLERRK	#: AAB46 11 GSDPLWDWNV NKTKTALGFL GVQSSGIMLT	21 TWNTSNPDFT LWIVCWADLF FWLVALVCAL	 KCFQNTVLVW YSFWERSRGI AILRSKIMTA	VPCFYLWACP FLAPVFLVSP LKEDAQVDLF	PFYFLYLSRH TLLGITTLLA RDITFYVYFS	120 180
	SEQ ID NO:22 O Protein Accession 1 MALRGFCSAD DRGYIOMTPL TFLIQLERRK LLLIQLVLSC	AAB46 11	616 21 TWNTSNPDFT LWIVCWADLF FWLVALVCAL TIHDPNFCPE	KCFQNTVLVW YSFWERSRGI AILRSKIMTA SSASFLSRIT	VPCFYLWACF FLAPVFLVSP LKEDAQVDLF FWWITGLIVR	PFYFLYLSRH TLLGITTLLA RDITFYVYFS GYRQPLEGSD	120 180 240
55	SEQ ID NO:22 O Protein Accession 1 MALRGFCSAD DRGYIQMIPL TPLIQLERRK LLLIQLVLSC LWSLNKEDTS	11 GSDPLWDWNV NKTKTALGFL GVQSSGIMLT FSDRSPLFSE EQVVPVLVKN	516 21 TWNTSNPDFT LMIVCWADLF FWLVALVCAL TIHDPNPCPE WKKECAKTRK	KCFQNTVLVW YSFWERSRGI AILRSKIMTA SSASFLSRIT QPVKVVYSSK	VPCFYLWACP FLAPVFLVSP LKEDAQVDLF FWWITGLIVR DPAQPKESSK	 PFYFLYLSRH TLLGITTLLA RDITFYVYFS GYRQPLEGSD VDANEEVEAL	120 180 240 300
	SEQ ID NO:22 Q Protein Accession I MALRGFCSAD DRGYIQMTPL TFLIQLERRK LLLIQLVLSC LWSINKEDTS IVKSPQKEWN	11	21 TWNTSNPDFT LWIVCWADLF FWLVALVCAL THDPNPCPE WKKECAKTRK FGPYFLMSFP	KCFQNTVLVW YSFWERSRGI AILRSKIMTA SSASFLSRIT QPVKVVYSSK FKAIHDLMMF	VPCFYLWACP FLAPVFLVSP LKEDAQVDLF FWWITGLIVR DPAQPKESSK SGPQILKLLI	PFYFLYLSRH TLLGITTLLA RDITFYVYFS GYRQPLEGSD VDANEEVEAL KFVNDTKAPD	120 180 240 300 360
55	SEQ ID NO:22 Q Protein Accession MALRGFCSAD DRGYIQMTPL TPLIQLERRK LLLIQLVLSC LWSLNKEDTS LVKSPQKEWN WQGYFYTVLL	11	21 TWNTSNPDFT LWIVCWADLF FWLVALVCAL TIHDPNPCPE WKKECAKTRK FGPYFLMSFF LEQYFHICFV	KCFQNTVLVW YSFWERSRGI AILRSKIMTA SSASFLSRIT QPVKVVYSSK PKAIHDLMMF SGMRIKTAVI	VPCFYLWACP FLAPVFLVSP LKEDAQVDLF PWWITGLIVR DPAQPKESSK SGPQILKLLI GAVYRKALVI	PFYFLYLSRH TLLGITTLLA RDITFYVYFS GYRQPLEGSD VDANEEVEAL KFVNDTKAPD TNSARKSSTV	120 180 240 300 360 420
55	SEQ ID NO:22 O Protein Accession 1	11 1 GSDPLWDWNV NKTKTALGFL GVQSSGIMLT FSDRSPLFSE EQVVPVLVKN PSLFKVLYKN PSLFKVLYKN PSLFKVLYKL AQRFMDLATY	21 TWNTSNPDFT LMIVCWADLF FWLVALVCAL TIHDPNPCPE WKKECAKTRK FGPYFLMSFF LEQYFHICFV INMIWSAPLQ	 KCFQNTVLVW YSFWERSRGI AILRSKIMTA SSASFLSRIT QPVKVVYSSK FKAIHDLMMF SGMRIKTAVI VILALYLLWL	VPCFYLWACF FLAPVFLVSP LKEDAQVDLF FWWITGLIVF DPAQPKESSK SGPQILKLLI GAVYRKALVI NLGPSVLAGV	PFYFLYLSRH TLLGITTLLA RDITFYVYFS GYRQPLEGSI VDANEEVEAL KFVNDTKAPD TNSARKSSTV AVMVLMVFVN	120 180 240 300 360 420 480
55 60	SEQ ID NO:22 Q Protein Accession 1	AAB46 11 GSDPLWDWNV NKTKTALGFL GVQSGIMLT FSDRSPLFSE EQVVPVLVKN PSLFKVLYKT FVTACLQTLU QVAHMKSKDN	21 TWNTSNPDFT LMIVCWADLF FWLVALVCAL THDPNPCPE WKKECAKTRK FGPYFLMSFF LHQYFHICFV INMIWSAPLQ RIKLMNEILM	KCFONTVLVW YSFWERSRGI ALLRSKIMTA SSASFLSRIT OPVKVVYSSK FKAIHDLMMF SGMRIKTAVI VILALYLLWL GIKVLKLYAW	VPCFYLWACF FLAPVFLVSP LKEDAQVDLF FWWITGLIVR DPAQPKESSK SGPQILKLLI GAVYRKALVI MLGPSVLAGV ELAFKDKVLA	PFYFLYLSRH TLLGITTLLX RDITFYVYFS GYRQPLEGSD VDANEEVEAL KFYNDTKAPD TNSARKSSTV AVMVLMVFVN IRQEELKVLK	120 180 240 300 360 420 480 540
55	SEQ ID NO:22 Q Protein Accession I	11	21 TWNTSNPDFT LMIVCWADLF FWLVALVCAL THDPNPCPE WKKECAKTRK FGPYPLMSFP LHQYFHICFV INMIWSAPLQ RIKLMNEILN ALCTFAVYVT	KCFQNTVLVW YSFWERSRGI AILRSKIMTA SSASFLSRIT QPVKVVYSSK FKAIHDLMMF SGMRIKTAVI VILALYLLWL GIKVLKLYAW IDENNILDAQ	VPCFYLWACF FLAPVFLVSP LKEDAQVOLF FWWITGLIVR DPAQPKESSK SGPQILKLLI GAVYRKALVI NLGPSVLAGV ELAFKDKVLA TAFVSLALFN	PFYFLYLSRH TLLGITTLLA RDITFYVYFS GYROPLEGSD VDANEEVEAL KFVNDTKAPD TNSARKSSTV AVMVLMYFVN IRQEELKVLK ILRFPLNILP	120 180 240 300 360 420 480
55 60	SEQ ID NO:22 Q Protein Accession I HALRGFCSAD DRGYIQMTPL TFLIQLERRK LLLIQLVLSC LWSLNKEDTS LWSLNKEDTS SIVSSPQKEWN WQGYFYTVLL GEIVNLMSVD AVMAMKTKTY KSAYLSAVGT MVISSIVQAS	AAB46 11 GSDPLWDWNV NKTKTALGFL GVQSGIMLT FSDRSPLFSE EQVVPVLVKN PSLFKVLYKT FVTACLQTLU QVAHMKSKDN	21 TWNTSNPDFT LMIVCWADLF FWLVALVCAL TIHDPNPCPE WKKECAKTRK FGPYPLMSFF LHQYFHICFV INMIWSAPLQ RIKLMNEILN ALCTFAVYVT SHEELEPDSI	KCFQNTVLW YSFWERSRGI AILRSKIMTA SSASFLSRIT QPVKVVYSSK FKAIHDLMMF SGMRIKTAVI VILALYLLWL GIKVLKLYAU IDENNILDAQ ERRFVKDGGG	VPCFYLWACE FLAPVFLVSP LKEDAQVDLF FWWITGLIVR DPAQPKESS GGPQILKLLI GAVYRKALVI NLGPSVLAGV ELAFKDKVLA TAFVSLALFN TNSITVRNAT	PFYFLYLSRH TLLGITTLLA RDITFYVYFS GYRQPLEGSD VDANEEVEAL KFYNDTKAPD TNSARKSSTV AVMVLMVFVN IRQEELKVLK LLRFPLNILP FTWARSDPPT	120 180 240 300 360 420 480 540 600
55 60	SEQ ID NO:22 Q Protein Accession 1	AAB46 11 GSDPLWDWNV NKTKTALGFL GVQSSGIMLT FSDRSPLFSE EQVVPVLVKN PSLFKVLYKT FVTACLQTLV AQRFMDLATY QVAHMKSKDN FTWVCTPFLV VSLKRLRIFL GALVAVVGQV QLEEPYYRSV	21 TWNTSNPDFT LMIVCWADLF FWLVALVCAL THDPNPCPE WKKECAKTRK FGPYFLMSFF LHQYFHICFV INMIWSAPLQ RIKLMNEILN ALCTFAVYVT SHEELEPDSI GCGKSSLLSA IQACALLPDL	KCFQNTVLVW YSFWERSRGI ALLRSKIMTA SSASFLSRIT QPVKVVYSSK FKAIHDLMMF FKAIHDLMMF GIKVLKLYLWL GIKVLKLYAW IDENNILDAQ ERRFVKDGGG LLAEMDKVEG EILPSGDRTE	VPCFYLWACF FLAPVFLVSP LKEDAQVDLF FWWITGLIVR DPAQPKESSK SGPQILKLLI GAVYRKALVI NLGFSVLAGV ELAFKDKVLA TAFVSLALFAT TNSITVRNAT TNSITVRNAT HVAIKGSVAY IGEKGVNLSG	PFYFLYLSRH TLLGITTLLA RDITFYVYFS GYRQPLEGSD VDANEEVEAL KFVNDTKAPD TNSARKSSTV AVMVLMVFVN IRQEELKVLK ILRFPLNILP FTWARSDPPT VPQQAWIQND GQKQRVSLAR	120 180 240 300 360 420 480 540 600 660
55 60	SEQ ID NO:22 Q Protein Accession I	11	21 TWNTSNPDFT LWIVCWADLF FWLVALVCAL TIHDPNPCPE WKKECAKTRK FGPYPHUSFF LHQYFHICFV INMIWSAPLQ RIKLMNEILN ALCTFAVYVT SHEELEPDSI GCGKSSLLSA LQACALLPDL LVGKHIFENV	KCFQNTVLW YSFWERSRGI AILRSKIMTA SSASFLSRIT QPVKVVYSSK FKAIHDLMMF SGMRIKTAVI VILALYLLWL GIKVLKLYAW IDENNILDAQ ERRFVKDGGG LLAEMDKVEG EILPSGDRTE IGPKGMLKNK	VPCFYLWACE FLAPVFLVSP LKEDAQVDLF FWWITGLIVR DPAQPKESS SGPQILKLLI GAVYRKALVI NLGPSVLAGV ELAFKDKVLA TAFVSLALFN TNSITVRNAT HVAIKGSVAY IGEKGVNLSG TRILVIHSMS	PFYFLYLSRH TLLGITTLLA RDITFYVYFS GYRQPLEGSD VDANEEVEAL KFYNDTKAPD TNSARKSSTV AVMVLMVPVN IRQEELKVLK ILRFPLNILP FTWARSDPPT VPQQAWIQND GQKQRVSLAR YLPQVDVIIV	120 180 240 300 360 420 480 540 660 720 780 840
55 60 65	SEQ ID NO:22 O Protein Accession 1	AB46 11	21 TWNTSNPDFT TWNTSNPDFT LMIVCWADLF FWLVALVCAL TIHDPNPCPE WKKECAKTRK FGPYPLMSFP LHQYFHICFV INMIWSAPLQ RIKLMNEILN ALCTFAVYVT SHEELEPDSI GCGKSSLLSA IQACALLPDL HVGKHIFENV AFAEFLRTYA	KCFQNTVLW YSFWERSRGI AILRSKIMTA SSASFLSRIT QPVKVVYSSK FKAIHDLMMF SGMRIKTAVI VILALYLLWL GIKVLKLYAW IDENNILDAQ ERRPVKDGGG LLAEMDKVEG EILPSGDRTE EIGPKGMLKNK STEQEQDAEE	VPCFYLWACE FLAPVFLVSP LKEDAQVDLF FWWITGLIVR DPAQPKESSK SGPQILKLLI GAVYRKALVI NLGPSVLAGV ELAFKDKVLA TAFVSLALFN TNSITVRNAT HVAIKGSVAY IGEKGVNLSG TRILVTHSMS NGVTGVSGPG	PFYFLYLSRH TLLGITTLLA RDITFYVYFS GYRQPLEGSD VDANEEVEAL KFVNDTKAPD TNSARKSSTV AVMVLMVFVN IRQEELKVLK ILRFPLNILP FTWARSDPPT VPQQAWIQND GQKQRVSLAR YLPQVDVIIV KEAKQMENGM	120 180 240 300 360 420 480 540 660 720 780 840 900
55 60	SEQ ID NO:22 OP Protein Accession I HALRGFCSAD DRGYIQMTPL TFLIQLERRK TFLIQLVISC LWSLNKEDTS IVKSPQKEWN WQGYFYTVLL GEIVNLMSVD AVMAMKTKTY KSAYLSAVGT MVISSIVQAS LNGITFSIPE SLRENILFGC AVYSNADIYL MSGGKISEMG LVTDSAGKQL	11 11 GSDPLWDWNV NKTKTALGFL GVQSSGIMLT FSDRSPLFSE EQVVPVLVKN PSLFKVLYKN PSLFKVLYKN AQRFMDLATY QVAHMKSKDN FTWVCTPFLV VSLKRLRIFL GALVAVVGQV QLEEPYYRSV FDDPLSAVDA SYQELLARDG QRQLSSSSSY	21 TWNTSNPDFT LMIVCWADLF FWLVALVCAL TIHDPNPCPE WKKECAKTRK FGFYFLMSFF LHQYFHICFV INMIWSAPLQ RIKLMNEILN ALCTFAVYUT SHEELEPDSI GCGKSSLLSA LQACALLPDL HVGKHIFENV AFAEFLRTYA SGDISRHHNS	KCFQNTVLVW YSFWERSRGI AILRSKIMTA SSASFLSRIT QPVKVVYSSK FKAIHDLMMF SGMRIKTAVI VILALYLLWL GIKVIKLYAW IDENNILDAG ERRPVKDGGG LLAEMDKVEG EILPSGDRTE IGPKGMLKNI STEQEQDAEE TAELQKAEAK	VPCFYLWACE FLAPVFLVSP LKEDAQVDLF FWWITGLIVR DPAQPKESSK SGPQILKLLI GAVYRKALVI NLGPSVLAGV ELAFKDKVLA TAFVSLALFN TNSITVRNAT HVAIKGSVAY IGEKGVNLSG TRILVTHSM NGVTGVSGPG KEETWKLMEA	PFYFLYLSRH TLLGITTLLA RDITFYVYFS GYRQPLEGSD VDANEEVEAL KFVNDTKAPD KFVNDTKAPD KFVNDTKAPD KFVNUTVVN IRQEELKVLK ILRFPLNILF FTWARSDPPT VPQQAWIQND GQXQRVSLAR YLPQVDVIIV KEAKQMENGM DKAQTGQVKL	120 180 240 300 360 420 480 540 660 720 780 840 900 960
55 60 65	SEQ ID NO:22 Q Protein Accession 1	11 GSDPLWDWNV NKTKTALGFL GVQSSGIMLT FSDRSPLFSE EQVVPVLVKN PSLFKVLYKT FVTACLQTLV AQRFMDLATY QVAHMKSKDN FTWVCTPFLV VSLKRLRIFL GALVAVVGQV QLEEPYYRSV FDDPLSAVDA SYQELLARDA GQRQLSSSSSY GLFISFLSIF	21 TWNTSNPDFT LMIVCWADLF FWLVALVCAL TIHDPNPCPE WKKECAKTRK FGPYFLMSFF LHQYFHICFV INMIWSAPLQ RIKLMNEILN ALCTFAVYVT SHEELEPDSI GCGKSSLLSA IQACALLPDL HVGKHIFENV AFAEFLRTYA SGDISRHHNS LFMCNHVSAL	KCFQNTVLVW YSFWERSRGI ALLRSKIMTA SSASFLSRIT QPVKVVYSSK FKAIHDLMMF SGMRIKTAVI VILALYLLWL GIKVLKLYAW IDENNILDAQ ERRPVKDGGG LLAEMDKVEG EILPSGDRTE IGPKGMLKNK STEQEQDAEE TABELQKAEAK ASNYWLSLWT	VPCFYLWACE FLAPVFLVSP LKEDAQVDLF FWWITGLIVR DPAQPKESSK SGPQILKLLI GAVYRKALVI NLGPSVLAGV ELAFKDKVLA TAFVSLALFIN TNSITVRNAT HVAIKGSVAY IGEKGVNLSG TRILVTHSMS NGVTGVSGPG KEBTWKLMEA DDPIVNGTQE	PFYFLYLSRH TLLGITTLLA RDITFYVYFS GYRQPLEGSD VDANEEVEAL KFVNDTKAPD TNSARKSSTV AVMVLMVFVN IRQEELKVLK ILRFPLNILP FTWARSDPPT VPQQAWIQND GQKQRVSLAR YLPQVDVIIV KEAKQMENGM DKAQTGQVKL HTKVRLSVYG	120 180 240 300 420 480 540 660 720 780 900 960 1020
55 60 65	SEQ ID NO:22 Q Protein Accession 1	11	21 TWNTSNPDFT LWIVCWADLF FWLVALVCAL TIHDPNPCPE WKKECAKTRK FGPYPLMSFP LHQYFHICFV INMIWSAPLQ RIKLMNELLM ALCTFAVYVT SHEELEPDSI GCGKSSLLSA IQACALLPDL HVGKHIFENV APAEFLRTYA SGDISRHHNS LPMCNHVSAL GILASRCLHV	KCFQNTVLW YSFWERSRGI AILRSKIMTA SSASFLSRIT QPVKVVYSSK FKAIHDLMMF SGMRIKTAVI VILALYLLWIL GIKVLKLYAW IDENNILDAQ ERRFVKDGGG LLAEMDKVEG EILPSGDRTE IGPKGMLKNK STEQEQDAEE TAELQKAEAK ASNYWLSLWT DLLHSILRSF	VPCFYLWACE FLAPVFLVSP LKEDAQVDLF FWWITGLIVR DPAQPKESS SGPQILKLLI GAVYRKALVI NLGPSVLAGV ELAFKDKVLA TAFVSLALFN TNSITVRNAT HVAIKGSVAY IGEKGVNLSG TRILVTHSMS NGVTGVSGPG KEBTWKLMEA DDPIVNGTQE MSFFERTPSG	PFYFLYLSRH TLLGITTLLA RDITFYVYFS GYRQPLEGSD VDANEEVEAL KFYNDTKAPD TNSARKSSTV AVMVLMVFVN IRQEELKVLK ILRFPLNILP FTWARSDPPT VPQQAWIQND GQKQRVSLAR YLPQVDVIIV KEAKQMENGM DKAQTGQVKL HTKVRLSVYG NLVNRFSKEL	120 180 240 360 420 480 540 660 720 780 840 900 1020 1080
55 60 65 70	SEQ ID NO:22 O Protein Accession 1	11	21 TWNTSNPDFT TWNTSNPDFT LMIVCWADLF FWLVALVCAL TIHDPNPCPE WKKECAKTRK FGPYPLMSFP LHQYFHICFV INMIWSAPLQ RIKLMNEILN ALCTFAVYVT SHEELEPDSI GCGKSSLLSA LQACALLPDL HVGKHIFENV AFAEFLRTYA SGDISRHINS LFMCNHVSAL LFMCNHV LF	KCFQNTVLW YSFWERSRGI AILRSKIMTA SSASFLSRIT QPVKVVYSSK FKAIHDLMMF SGMRIKTAVI VILALYLLWL GIKVLKLYAW IDENNILDAQ ERRPVKDGGG LLAEMDKVEG EILPSGDRTE EIGPKGMLKNK STEQEQDAEE TAELQKAEAK ASNYWLSLWT DLHSILRSP ATPIAAIIIP	VPCFYLWACE FLAPVFLVSP LKEDAQVDLF FWWITGLIVR DPAQPKESSK SGPQILKLLI GAVYRKALVI NLGPSVLAGV ELAFKDKVLA TAFVSLALFN TNSITVRNAT HVAIKGSVAY IGEKGVNLSG TRILVTHSMS NGVTGVSGPG KEBTWKLMEA DDPIVNGTQE MSFFERTPSG PLGLIYFFVQ	PFYFLYLSRH TLLGITTLLA RDITFYVYFS GYRQPLEGSD VDANEEVEAL KFVNDTKAPD TNSARKSSTV AVMVLMVFVN IRQEELKVLK ILRFPLNILP FTWARSDPPT VPQQAWIQND GQKQRVSLAR YLPGVDVIIV KEAKQMENGM DKAQTGQVKL HTKVRLSVYG NLVNRFSKEL RFYVASSRQL	120 180 240 360 420 480 540 660 720 780 840 900 960 1020 1080 1140
55 60 65 70	SEQ ID NO:22 O Protein Accession 1 MALRGFCSAD DRGYIQMTPL TFLIQLERRK LLLIQLVLSC LWSLMKEDTS IVKSFQKEWN WQGYFYTVLI GEIVNILMSVD AVMAMKTKTY KSAYLSAVGT MVISSIVQAS LNGITFSIPE SLRENILFGC AVYSNADIYL MSGGKISEMG LVTTDSAGKQL SVYWDYMKAI ALGISQGIAV DTVDSMIPEV KRLESVSRSP	1#: AAB46 11 GSDPLWDWNV NKTKTALGFL GVQSSGIMLT FSDRSPLFSE EQVVPVLVKN PSLFKVLYKT FVTACLQTLV AQRFMDLATY QVAHMKSKDN FTWVCTPFLV VSLKRLRIFL GALVAVVGQV QLEEPYYRSV FDDPLSAVDA GYQLLARDG QRQLSSSSY GLFISFLSIF FGYSMAVSIG KMFMGSLFN VYSHFNETLL	21 TWNTSNPDFT LMIVCWADLF FWLVALVCAL TIHDPNPCPE WKKECAKTRK FGFYFLMSFF LHQYFHICFV INMIWSAPLQ RIKLMNEILN ALCTFAVYVI SHEELEPDSI GCGKSSLLSA LQACALLPDL HVGKHIFENV AFAEFLRTYA SGDISRHHNS LPMCNHVSAL GILASRCLHV UIGACIVILL GVSVIRAFEE	KCFQNTVLVW YSFWERSRGI AILRSKIMTA SSASFLSRIT QPVKVVYSSK FKAIHDLMMF SGMRIKTAVI VILALYLLWL GIKVIKLYAW IDENNILDAG ERRPVKDGGG LLAEMDKVEG EILPSGDRTE IGPKGMLKNK STEQEQDAEE TAELQKAEAK ASNYWLSLWT DLLHSILRSP ATPIAAIIIP QERFIHQSDL	VPCFYLWACE FLAPVFLVSP LKEDAQVDLF FWWITGLIVR DPAQPKESSK SGPQILKLLI GAVYRKALVI NLGPSVLAGV ELAFKDKVLA TAFVSLALFN TMSITVRNAT HVAIKGSVAY IGEKGVNLSG TRILVTHSM NGVTGVSGPG KEETWKLMEA DDPIVNGTQE MSFFERTPSQ FLGLIYFFVQ KVDENQKAYY	PFYFLYLSRH TLIGITTLLA RDITFYVYFS GYRQPLEGSD VDANEEVEAL KFVNDTKAPD TINSARKSSTV AVMVLMVFVN IRQEELKVLK ILRFPLNILF FTWARSDPPT VPQQAWIQND GQXQRVSLAR YLPQVDVIIV KEAKQMENGM DKAQTGQVKL HTKVRLSVYG NLVNRFSKEL RFYVASSRQL PSIVANRWLA	120 180 240 300 360 420 480 660 720 780 960 1020 1020 1140 1200
55 60 65	SEQ ID NO:22 Q Protein Accession 1	11 GSDPLWDWNV NKTKTALGFL GVQSSGIMLT FSDRSPLFSE EQVVPVLVKN PSLFKVLYKT FVTACLQTLV AQRFMDLATY QVAHMKSKDN FTWVCTPFLV VSLKRLRIFL GALVAVVGQV QLEEPYYRSV FDDPLSAVDA SYQELLARDG QRQLSSSSSY GLFISFLSIF FGYSMAVSIG IKMFMGSLFN VYSHFNETLL VLFAALFAVI	21 TWNTSNPDFT LMIVCWADLF FWLVALVCAL TIHDPNPCPE WKKECAKTRK FGFYFLMSFF LHQYFHICFV INMIWSAPLQ RIKLMNEILN ALCTFAVYVT SHEELEPDSI GCGKSSLLSA IQACALLPDL HVGKHIFENV AFAEFLRTYA SGDISRHHNS LFMCNHVSAL GILASRCLHV VIGACIVILI GVSVIRAFEE SRHSLSAGLV	KCFQNTVLW YSFWERSRGI AILRSKIMTA SSASFLSRIT QPVKVVYSSK FKAIHDLMMF SGMRIKTAVI VILALYLLWIL GIKVLKLYAW IDENNILDAQ ERRPVKDGGG LLAEMDKVEG EILPSGDRTE IGPKGMLKNK STEQEQDAEE TAELQKAEAK ASNYWLSLWT DLLHSILRSP ATPIAAIIIP QERFIHQSDL GLSVSYSLQV	VPCFYLWACE FLAPVFLVSP LKEDAQVDLF FWHITGLIVR DPAQPKESS SGPQILKLLI GAVYRKALVI NLGPSVLAGV ELAFKDKVLA TAFVSLALFN TNSITVRNAT HVAIKGSVAJ IGEKGVNLSG TRILVTHSMS NGVTGVSGPG KEETWKLMEA DDPIVNGTQE MSFFERTPSG PLGLIYFFVQ KVDENQKAYY TTYLNWLVRM	PFYFLYLSRH TLLGITTLLA RDITFYVYFS GYRQPLEGSD VDANEEVEAL KFYNDTKAPD TNSARKSSTV AVMVLMVPVN IRQEELKVLK ILRFPLNILP FTWARSDPPT VPQQAWIQND GQXQRVSLAR YLPQVDVIIV KEAKQMENGM DKAQTGQVKL HTKVRLSVYG NLVNRFSKEL RFYVASSRQL FSIVANRWLA SSEMETMIVA	120 180 240 360 420 480 540 660 720 780 840 900 960 1020 1080 1140
55 60 65 70	SEQ ID NO:22 O Protein Accession 1	11 11 CSDPLWDWNV NKTKTALGFL GVQSSGIMLT FSDRSPLFSE EQVVPVLVKN PSLFKVLYKT FVTACLQTLV AQRFMDLATY QVAHMKSKDN FTWVCTPFLV VSLKRLRIFL GALVAVVGQV QLEEPYYRSV FDDPLSAVDA SYQELLARDG QRQLSSSSY GLFISFLSIF FGYSMAVSIG IKMFMGSLFN VYSHFNETLL VLFAALFAVI EKEAPFWQIQE AGKSSLTLGL	21 TWNTSNPDFT LMIVCWADLE FWLVALVCAL TIHDPNPCPE WKKECAKTRK FGFYFLMSFF LHQYFHICFV INMIWSAPLQ RIKLMNEILN ALCTFAVYVT SHEELEPDSI GCGKSSLLSA LQACALLPDL HVGKHIPENV AFAEFLRTYA SGDISRHINS LFMCNHVSAL GULASRCLHV UIGACIVILL GVSVIRAFEE SRHSLSAGLV TAPPSSWPQV FRINESAEGE	KCFQNTVLVW YSFWERSRGI AILRSKIMTA SSASFLSRIT QPVKVVYSSK FKAIHDLMMF SGMRIKTAVI VILALYLLWL GIKVLKLYAW IDENNILDAQ ERRPVKDGGG LLAEMDKVEG EILPSGDRTE EIGPKGMLKNK STEQEQDAEE TAELQKAEAK ASNYWLSLWT DLHSILRSP ATPIAAIIIP QERFIHQSDL GLSVSYSLQV GRVEFRNYCL IIIDGINIAK	VPCFYLWACE FLAPVFLVSP LKEDAQVDLF FWWITGLIVR DPAQPKESSK SGPQILKLLI GAVYRKALVI NLGPSVLAGV ELAFKDKVLA TAFVSLALFN TNSITVRNAT HVAIKGSVAY IGEKGVNLSG KEGTWKLMEA DDPIVNGTQE MSFFERTPSG PLGLIYFFVQ KVDENQKAYY TTYLNWLVRM TYLNWLVRM TYLNWLVRM TYLNWLVRK IGLHDLFYKI IGLHDLFYKI	PFYFLYLSRH THEGITTLLA RDITFYVYFS GYRQPLEGSD VDANEEVEAL KPVNDTKAPD TNSARKSSTV AVMVLMVPVN IRQEELKVLK IRQEELKVLK IRQEELKVLK IRQEELKVLK IRQEELKVLK IRQEELKVLK IRQEELKVLK IRQEELKVLK IRQEELKVLK IRGEELKVLK IRQEELKVLK IRQEELKVLK IRQEELKVLK IRQEELKVLK IRQEELKVLK VPQQAWIQND GQKQRVSLAR VLPQVDVIIV KEAKQMENGM DKAQTGQVKL HTKVRLSVYG NLVNRFSKEL RFYVASSRQL PSIVANRWLA SSEMETNIVA SSEMETNIVA SHINVTINGG TIIPQDFVLF	120 180 240 300 360 480 540 660 720 780 840 960 1020 1080 1140 1200 1260
55 60 65 70	SEQ ID NO:22 Q Protein Accession 1 MALRGFCSAD DRGYIQMTPL TFLIQLERRK LLLIQLVLSC LWSLMKEDTS IVKSFQKEWN WQGYFYTVLI AVMAMKTKTY KSAYLSAVGT MVISSIVQAS LNGTIFSIPE SLRENILFGC AVYSNADIYL MSGGKISEMG LVTDSAGKQL SVYWDYMKAI ALGISQGIAV DTVDSMIPEV KRLESVSRSP VRLECVGNCI VERLKEYSET EKVGIVGRTG SGSLRMNLDP	1#: AAB46 11 GSDPLWDWNV NKTKTALGFL GVQSSGIMLT FSDRSPLFSE EQVVPVLVKN PSLFKVLYKT FVTACLQTLV AQRFMDLATY QVAHMKSKDN FTWVCTPFLV VSLKRLRIFL GALVAVVGQV QLEEPYYRSV FDDPLSAVDA GVQLSSSSY GLFISFLSIF FGYSMAVSIG KMFMGSLFN VYSHFNETLL VLFAALFAVI EKEAPWQIQE EKEAPWQIQE AGKSSLTLGL FSQYSDEEWW	21 TWNTSNPDFT LMIVCWADLF FWLVALVCAL TIHDPNPCPE WKKECAKTRK FGFYFLMSFF LRQYFHICFV INMIWSAPLQ RIKLMNEILN ALCTFAVYVI SHEELEPDSI GCGKSSLLSA LQACALLPDL HVGKHIFENV SGDISRHHNS LFMCNHVSAL GILASRCLHV GUSVIRAFEE SRHSLSAGLV TAPPSSWPQU FRINESAEGE TSLELAHLKD	KCFQNTVLVW YSFWERSRGI AILRSKIMTA SSASFLSRIT QPVKVVYSSK FKAIHDLMMF SGMRIKTAVI VILALYLLWL GIKVIKLYAW IDENNILDAG ERRPVKDGGG LLAEMDKVEG EILPSGDRTE IGPKGMLKNK STEQEQDAEE TAELQKAEAK ASNYWLSLWT DLHSILRST QERFIHQSDL GLSVSYSLQV GRVEFRNYCL IIIDGINIAK FVSALPDKLD	VPCFYLWACE PLAPVFLVSP LKEDAQVDLF FWWITGLIVR DPAQPKESSK SGPQILKLLI GAVYRKALVI NLGPSVLAGV ELAFKDKVLA TAFVSLALFN TMSITVRNAT HVAIKGSVAY IGEKGVNLSG REILVTHSMS NGVTGVSGPG KEETWKLMEA DDPIVNGTQE MSFFERTPSC KVDENQKAYY TTYLNWLVRM RYREDLDFVLI IGLHDLRFKI HECAEGGENL	PFYFLYLSRH PFYFLYLSRH TLLGITTLLA RDITFYVYFS GYRQPLEGSD VDANEEVEAL KFVNDTKAPD TNSARKSSTV AVMVLMVPVN IRQEELKVLK ILRFPLNILP FTWARSDPPT VPQQAWIQND GQKQRVSLAR YLPQVDVIIV KEAKQMENGM DKAQTGQVKL HTKVRLSVYG NLVNRFFSKEL FSIVANRWLA SSEMETNIVA RHINVTINGG TIIPQDPVLP SVGQRQLVCL	120 180 240 300 360 420 480 660 720 840 900 900 1020 1140 1200 1260 1320
55 60 65 70	SEQ ID NO:22 Q Protein Accession 1	11 11 CSDPLWDWNV NKTKTALGFL GVQSSGIMLT FSDRSPLFSE EQVVPVLVKN PSLFKVLYKT FVTACLQTLV AQRFMDLATY QVAHMKSKDN FTWVCTPFLV VSLKRLRIFL GALVAVVGQV QLEEPYYRSV FDDPLSAVDA SYQELLARDG QRQLSSSSY GLFISFLSIF FGYSMAVSIG IKMFMGSLFN VYSHFNETLL VLFAALFAVI EKEAPFWQIQE AGKSSLTLGL	21 TWNTSNPDFT LWIVCWADLF FWLVALVCAL TIHDPNPCPE WKKECAKTRK FGPYPLMSFF LHEQYFHICFV INMIWSAPLQ RIKLMNEILN ALCTFAVYVT SHEELEPDSI GCGKSSLLSA IQACALLPDL HVGKHIFENV AFAEFLRTYA SGDISRHHNS GILASRCLHV VIGACIVILL GVSVIRAFEE SRHSLSAGLV TAPPSSWPQV FRINESAEGE TSLELAHLKD DLETDDLIQS	KCFQNTVLW YSFWERSRGI AILRSKIMTA SSASFLSRIT QPVKVVYSSK FKAIHDLMMF SGMRIKTAVI VILALYLLWIL GIKVLKLYAW IDENNILDAQ ERRPVKDGGG LLAEMDKVEG EILPSGDRTE IGPKGMLKNK STEQEQDAEE TAELQKAEAK ASNYWLSLWT DLLHSILRSP ATPIAAIIIP QERFIHQSDL GLSVSYSLQV GRVEFRNYCL IIIDGINIAK FVSALPDKLD TIRTQFEDCT	VPCFYLWACE PLAPVFLVSP LKEDAQVDLF FWWITGLIVR DPAQPKESSK SGPQILKLLI GAVYRKALVI NLGPSVLAGV ELAFKDKVLA TAFVSLALFN TMSITVRNAT HVAIKGSVAY IGEKGVNLSG REILVTHSMS NGVTGVSGPG KEETWKLMEA DDPIVNGTQE MSFFERTPSC KVDENQKAYY TTYLNWLVRM RYREDLDFVLI IGLHDLRFKI HECAEGGENL	PFYFLYLSRH PFYFLYLSRH TLLGITTLLA RDITFYVYFS GYRQPLEGSD VDANEEVEAL KFVNDTKAPD TNSARKSSTV AVMVLMVPVN IRQEELKVLK ILRFPLNILP FTWARSDPPT VPQQAWIQND GQKQRVSLAR YLPQVDVIIV KEAKQMENGM DKAQTGQVKL HTKVRLSVYG NLVNRFFSKEL FSIVANRWLA SSEMETNIVA RHINVTINGG TIIPQDPVLP SVGQRQLVCL	120 180 240 360 420 480 540 660 720 840 900 960 1020 1140 1200 1320 1380

SEQ ID NO:23 PAA2 DNA SEQUENCE

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	1 AGGAATCTGC ATCGGGCAGA CATGAGGATT CAGGATCATC CGAGAAGACG AGCCCACTGC	25-874 11 GCTCGGGTTC GGTCTCACAG CTGCAGTTAA AAGGGGTTCG CGGCTACTCT CTCAAGCCCC	(underlined seque 21 CGCAGATGCA CAGCCAAGGA TCCTGCTTGC AGTGCAAGCC GTGGGGCGAC GCTACATAGT	31 GAGGTTGAGG ACCTGGGAACA TCACTCCCAG GCTCATCGCC TCACCTGGGG	41 TGGCTGCGGG CGCTTCTCCC GGGCTTGTAG CCCTGGCAGG CCCAGATGGC CAGCACAACC	51 ACTGGAAGTC CCCTCCAGGC GGGGAGAGAC CAGCCCTGTT TCCTGACAGC TCCAGAAGGA	60 120 180 240 300 360
	1 AGGAATCTGC ATCGGGCAGA CATGAGGATCATC CGAGAAGACG AGCCCACTGC GGAGGGCTGT	26-874 11 GCTCGGGTTC GGTCTCACAG CTGCAGTTAA AAGGGGTTCG CGCTACTCT CTCAAGCCCC GAGCAGACCC	21	31 GAGGTTGAGG ACCTGGGGCC TCTGGCAACA TCACTCCAG GCTCATCGCC TCACCTGGGG TGAGTCCTTC	41 TGGCTGCGGG CGCTCCTCCC GGGCTTGTAG CCCTGGCAGG CCCAGATGGC CAGCACAACC CCCCACCCCG	51 ACTGGAAGTC CCCTCCAGGC GGGGAGAGAC CAGCCCTGTT TCCTGACAGC TCCAGAAGGA GCTTCAACAA	60 120 180 240 300 360 420
	1 AGGAATCTGC ATCGGGCAGA CATGAGGATT CAGGATCATC CGAGAAGACG AGCCCACTGC GGAGGGCTGT CAGCCTCCCC	26-874 11 GCTCGGGTTC GGTCTCACAG CTGCAGTTAA AAGGGGTTCG CGGCTACTCT CTCAAGCCCC CAGCAGACCC AACAAAGACC	21 CGCAGATGCA CAGCCAAGGCA AGTGCAAGCC GTGGGCGAC GCTACATAGT GGACAGCCAC ACCGCAATGA ACCGCAATGA	31 GAGGTTGAGG ACCTGGGACA TCACTCCCAG GCTCATCGCC TCACCTGGGG TGAGTCCTTC CATCATGCTG	41 TGGCTGCGGG CGCTCCTCCC GGGCTTGTAG CCCTGGCAGG CCAGATGGC CAGCACCCC CCCCACCCCG GTGAAGATGG	51 ACTGGAAGTC CCCTCCAGGC GGGGAGAGA CAGCCCTGTT TCCTGACAGG GCTTCAACAA GCTTCAACAA CATCGCCAGT	60 120 180 240 300 360 420 480
60	1 AGGAATCTGC ATCGGGCAGA CATGAGGATCATC CGAGAAGACG AGCCCACTGC GGAGGGCTGC CAGCCTCCCC CTCCATCACC	26-874 11 GCTCGGGTTC GGTCTCACAG CTGCAGTTAA AAGGGGTTCG CGGCTACTCT CTCAAGCCCC GAGCAGACCC AACAAAGACC TGGGCTGTCC	(underlined seque 21 CGCAGATGCA CAGCCAAGGA TCCTGCTTGC AGTGCAAGCC GTGGGGCGAC GCTACATAGT GGACAGCCA ACCGCAATGA GACCCCTCAC	31 GAGGTTGAGG ACCTGGGAACA TCACTCCAG GCTCATCGCC TCACCTGGGG TGAGTCCTTC CATCATGCTG CCTCTCCTCA	41 TGGCTGCGGG CGCTCCTCCC GGGCTTGTAG CCCTGGCAGGCCAGCCC CCCAGCACC CCCACCCC GTGAAGATGG CGCTGTGTCA	51 ACTGGAAGTC CCCTCCAGGC GGGGAGAGAC CAGCCCTGTT TCCTGACAGC TCCAGAAGGA GCTTCAACAA CATCGCCAGT CATCGCCAGT	60 120 180 240 300 360 420 480 540
	1 AGGAATCTGC ATCGGGCAGA CATGAGGATCATC CGAGAAGACG AGCCCACTGC GGAGGGCTGT CAGCCTCCCC CTCCATCACC CAGCTGCCTCC	26-874 11 GCTCGGGTTC GGTCTCACAG CTGCAGTTAA AAGGGGTTCG CGGCTACTCT CTCAAGCCCC GAGCAGACCC AACAAAGAC TGGGCTGTGC AATTCCGGCT	(underlined seque 21 CGCAGATGCA CAGCCAAGGA TCCTGCTTGC GTGGGGCGAC GCTACATAGT GGACAGCCAC ACCGCAATGA GACCCCTCAC GGGCCACCAC	31 GAGGTTGAGG ACCTGGGACA TCACTCCAG GCTCATCGCC TCACCTGGGG TGAGTCCTTC CATCATCCTC CCTCTCCTCA GTCCAGCCCC	41 TGGCTGCGGG CGCTCCTCCC GGGCTTGTAG CCCTGGCAGG CCCACCCCG CTGAAGATGG CGCTGTGTCA CAGTTACGCC	51 ACTGGAAGTC CCCTCCAGGC GGGGAGAGAC CAGCCCTGTT TCCTGACAGC TCCAGAAGGA GCTTCAACAA CATCGCCAGT TGCTGGCAC TGCCTCACAC	60 120 180 240 300 360 420 480 540 600
60	1 AGGAATCTGC ATCGGGCAGA CATGAGGATCATC CGAGAACAC GGAGGCCTGT CAGCCTCCCC CTCCATCACC CTCCATCACC CTTCGCGATGC	26-874 11 GCTCGGGTTC GGTCTCACAG CTGCAGTTAA AAGGGGTTCG CGGCTACTCT CTCAAGCCCC GAGCAGACCC AACAAAGACC AACAAAGACC ATTTCCGGCTGC ATTTCCGGCT	21 CGCAGATGCA CAGCCAAGGA CAGCCAAGGA TCCTGCTTGC AGTGCAAGCC GCTACATAGT GGACAGCCAC ACCGCAATGA ACCGCAATGA GACCCCTCAC GGGGCAGCAC CCATCATTGA	31 GAGGTTGAGG ACCTGGGGCC TCAGCTCCAG GCTCATCGCC TCACCTGGGG TGAGTCCTTC CATCATGCTG CCTCTCCTCA GTCCAGCCCC GCACCAGAAG	41 TGGCTGCGGG CGCTCCTCCC GGGCTTGTAG CCCTGGCAGG CCCAGATGGC CAGCACACCC GTGAAGATGG CGCTGTGTCA CAGTTACGCC TGTGAGAACC	51 ACTGGAAGTC CCCTCCAGGC GGGGAGAC CAGCCCTGTT TCCTGACAGC TCCAGAAGGA GCTTCAACAA CATCGCCAGT CTGCTGCAC TGCCTCACAC CCTACCCCGG	60 120 180 240 300 360 420 540 600 660
60	1 AGGAATCTGC ATCGGGCAGA CATGAGGATT CAGGATCATC CGAGAGGCTGT CAGCCTCCC CTCCATCAC CAGCTGCCTCCATCACC CAGCTGCCTCCATCACAC CAACATCACA CAACATCACA	26-874 11 GCTCGGGTTC GGTCTCACAG CTGCAGTTAA AAGGGGTTCG CGGCTACTCT CTCAAGCCCC CAGCAAGACCC AACAAAGACC TGGGCTGTGC ATTTCCGGCT GCCAACATCA GACACATCA GACACCATCG	21 CGCAGATGCA ACCCAAGATGCA ACCCAAGATGCA ACTGCAAGCC ACTGCAAGCC GTGGGCGAC GCTACATAGT GGACAGCCAC ACCGCAATGA GACCCTCAC GGGGCAGCA CCATCATTGA TGTGTGCCAG	31 GAGGTTGAGG ACCTGGGGCC TCATCCCAG GCTCATCGCC TCACCTGGGG TGAGTCCTTC CATCATGCTG GTCCAGCCCC GCACCAGGAG CGTGCAGGAAG	41 I TGGCTGCGGG CGGTCTCCC GGGCTTGTAG CCCTGGCAGG CCCAGCAGACC CAGCACACCCG GTGAAGATGG CGCTGTGTCA CAGTTACGCC TGTGAGAACG GGGGGCAAGG	51 ACTGGAAGTC CCCTCCAGGC GGGGAGAGAC CAGCCCTGTT TCCTGACAGC TCCAGAAGGA CATCGCCAGT CTGCTGGCAC TGCCTCACAC CCTACCCCGG ACTCCTGCCA	60 120 180 240 360 420 480 540 600 660 720
60 65	1 AGGAATCTGC ATCGGGCAGA CATGAGGATT CAGGATCATC CGAGAAGACG AGCCCACTGC GGAGGGCTGC CACCATCACC CTCCATCACC CTTCCATCACC CTTTCGGATGC CAACATCACA GGGTGACTCC GGGTGACTCC	26-874 11 GCTCGGGTTC GGTCTCACAG CTGCAGTTAA AAGGGGTTCG CGGCTACTCT CTCAAGCCCC GAGCAGACCC AACAAAGACC TGGGCTGTGC ATTTCCGGCT GCCAACATCA GACACCATGG GGGGGCCCTC	21 CGCAGATGCA CAGCCAAGGA TCCTGCTTGC AGTGCAAGCC GTGGGCGAC GCTACATAGT ACCGCAATGA ACCGCAATGA GACCCTCAC GGGCAGCAC CCATCATTGA TGTGTGCCAG TGGTCTCTAA	31 GAGGTTGAGG ACCTGGGAACA TCACTCCCAG GCTCATCGCC TCACCTGGGG TGAGTCCTTC CCTCTCCTCA GTCCAGCCCC GCACCAGAGC GCGTGCAGGAA CCAGTCTCTT	41 TGGCTGCGGG CGCTCCTCCC GGGCTTGTAG CCCTGGCAGC CCCAGCACC CCCACCCCC GTGAAGATGG CGCTGTGTCA CAGTTACGCC TGTGAGAAC CGGGGCAAGG CAAGGCATTA	51 ACTGGAAGTC CCCTCCAGGC GGGGAGAGAC CAGCCCTGTT TCCTGACAGC TCCAGAAGGA GCTTCAACAA CATCGCCAGT CTGCTGCCAC TGCTTACCCAG CCTTACCCCGG CCTACCCGG CTTCCTGCGGG TCTCCTGGGGG TCTCCTGGGGG	60 120 180 240 300 360 420 540 600 660
60	1 AGGAATCTGC ATCGGGCAGA CATGAGGATCATC CGAGAAGACG AGCCCACTGC GGAGGCCTGC CTCCATCACC CTCCATCACC CTTGCGATGAC CTTGCGATGC CAACATCAC CAACATCAC CCAGGATCCC CCAGGATCCC CCAGGATCCC	26-874 11 GCTCGGGTTC GGTCTCACAG CTGCAGTTAA AAGGGGTTCG CAGCAGACCC CAGCAGACCC TGGGCTGTGC ATTTCCGGCT GCCAACATCA GACACCATCG GGGGGCCTC TGTGCGATCA	(underlined seque 21 CGCAGATGCA CAGCCAAGGA TCCTGCTTGC GTGGGGCGAC GCTACATAGT GGACAGCCA ACCGCAATGA GACCCCTCAC GGGCAGCA TGTGTGCCAC TGTGTGCCAC TGTGTGCCAC TGTTGCCAC TGGTCTGTAA CCCGAAAGCC	31 GAGGTTGAGG ACCTGGGGCC TCATCCCAG GCTCATCGCC TCACCTGGGG TGAGTCCTTC CATCATGCTG GTCCAGCCCC GCACCAGGAG CGTGCAGGAAG	41 TGGCTGCGGG CGCTCCTCCC GGGCTTGTAG CCCTGGCAGC CCCACCCCG GTGAAGATGC CGCTGTGTCA CAGTTACGCC TGTGAGAACG CGGGGCAAGG CAAGGCATTA ACGAAAGTCT	51 ACTGGAAGTC CCCTCCAGGC GGGGAGAGAC CAGCCCTGTT TCCTGACAGC TCCAGAAGGA GCTTCAACAA CATCGCCAGT TGCTGGCAC TGCTCACAC CCTACCCGG ACTCCTGCGG ACTCCTGCGG GCAAATATGT	60 120 180 240 300 360 420 480 540 600 720 780
60 65	1 AGGAATCTGC ATCGGGCAGA CATGAGGATCAT CAGGATCATC CGAGAGACG AGCCTCCC CTCCATCACC CTCCATCACC CTGCGTGCCTC CTTGCGATGC CAACATCACA GGGTGACTCC CCAGGGTCACTC CCAGGGTCACTC CCAGGATCCG GGACTCGGATC GGGACTGGATC	26-874 11 GCTCGGGTTC GGTCTCACAG CTGCAGTTAA AAGGGGTTCG CGGCTACTCT CTCAAGCCC CAACAAAGACC AACAAAGACC AACAAAGACC GCCAACATCA GACACATCA GACACATCA GACACATCA CAGGGGCCTTC CTGTGCGATCA CAGGAGACGA	(underlined seque 21 CGCAGATGCA CAGCCAAGGA TCCTGCTTGC AGTGCAAGCC GTGGGCGAC GCTACATAGT GGACAGCCAC ACCGCAATGA ACCGCTACT ACCGCAATGA TGTGTGCCAG TGTGTGCCAG TGTGTGCCAG TGTGTCTGTAC TGTAGTGCAA	31 GAGGTTGAGG ACCTGGGACA TCACTCCAG GCTCATCGCC TCACCTGGGG TGAGTCCTTC CATCATGCTG CCTCTCCTCA GTCCAGGCCC GCACCAGAAG CGTGCAGGAA CCAGTCTCTT TGGTGTCTAC	41 TGGCTGCGGG CGCTCCTCCC GGGCTTGTAG CCCTGGCAGG CCCACCCCG GTGAAGATGG CGCTGTTACGCC TGTGAGATGC CGTGAGACG CAGTTACGCC TGTGAGAACC CGGGGCAAGG CAAGGCATTA ACGAAAGTCT CCCACCCACC	51 ACTGGAAGTC CCCTCCAGGC GGGGAGAGAC CAGCCCTGTT TCCTGACAGC TCCAGAAGGA GCTTCAACAA CATCGCCAGT TGCCTCACAC CCTACCCCGG ACTCCTCGCCA TCTCCTGGCG GCAAATATGT ACAGCCCATC	60 120 180 240 300 420 480 540 600 720 780 840 900 960
60 65	1 AGGAATCTGC ATCGGGCAGA CATGAGGATT CAGGATCATC CGAGAAGACG AGCCCACTGC GGAGGCCTCC CTCCATCACC CAGCTGCCTCCATCACC CAGCTGCCTC CAACACACAC GGATGACTCC CCAGGATCC CAACACC CCAGACC CCACACC CCAGACC CCAGACC CCACACC C	26-874 11 GCTCGGGTTC GGTCTCACAG CTGCAGTTAA AAGGGGTTCG CGGCTACTCT CTCAAGCCCC CAGCAGACCC AACAAAGACC TGGGCTGTGC ATTTCCGGCT GCCAACATCA GCAACATCA GACACCATGG GGGGGCCCTC TGTGGGATCA TCCACTTGGT TACGAACATT	(underlined seque 21 CGCAGATGCA CAGCCAAGGA TCCTGCTTGC GTGGGCGAC GCTACATAGT GCACACCAC GCGCAATGA GACCCCTCAC GGGGCAGCAC CCATCATTGA TGTGTGCCAG TGGTCTGTAA CCCGAAAGCC TGGAAAGCC TGGAAAACCC TGCAAGAACCC TTTGGGTCCC CTTTGGGCCT	31 GAGGTTGAGG ACCTGGGAACA TCACTCCCAG GCTCATCGCC TCACCTGGGG TCACCTGCGG TCACCTGCTC CCATCATCTC CCTCACCCC CCACCAGAG CCTCACCCC CCACCAGAG CCAGCAGGAA CCAGTCTTT TGGTGTCTAC TTAGACTGGT TGTTCACTCT CCTGGACTAC	41 TGGCTGCGGG CGCTCCTCCC GGGCTTGTAG CCCTGGCAGCCCCCCCCCC	51 ACTGGAAGTC CCCTCCAGGC GGGGAGAGAC CAGCCCTGTT TCCTGACAGC TCCAGACAGC CATCCCAGT CTCCTGCCAC TGCTCACAC TGCTCACAC TGCTCACAC TCCTGCCAC TCTCTGCGCA TCTCTTGCGG GCAAATATGT ACAGCCCATC AACCCTAAGC GTCACTAAGC GTCACTTAAT	60 120 180 240 300 360 420 480 540 600 720 780 840 900 900 1020
60 65	1 AGGAATCTGC ATCGGGCAGA CATGAGGATT CAGGATCATC CGAGAAGACG AGCCCACTGC GGAGGCTGT CAGCTCCCC CTCCATCACC CAGCTGCCTC CTTGCGATGAC CAACATCACA GGGTGACTCC CCAGGATCCG GGACTGGATCC CCAGGATCCG GGACTGGATC CCAGGATCCG ACCCTCCATT ACCCTCCATT AATCAACCTG	26-874 11 GCTCGGGTTC GGTCTCACAG CTGCAGTTAC CTGAGTTAC CTGAGTACTC CTCAAGCCC CAGCAGACCC TGGGCTGTGC ATTTCCGGCT GCCAACATCA GACACATGA GGGGGCCTC TGTGCGATCA CAGGAGACA TCACACTTGGT TACGAACATT GGGTTCGAAA	(underlined seque 21 CGCAGATGCA CAGCCAAGGA TCCTGCTTGC GGGCGAC GCTACATAGT GGACACCCACA GCGCAATGA GACCCCTCAC GGGCAGCAC CCATCATTGA TGTGTGCCAG TGGTCTGTAA CCCGAAAGCC TGAAGAACAA GTTTGGTCCC TCTTGGGCCT TCAGTGAGAC TCAGTGAGAC	31 GAGGTTGAGG ACCTGGGAACA TCACTCCCAG GCTCATCGCC TCACCTGGGG TGAGTCCTTC CATCATGCTG CCTCACAGAGG CCTCACAGAAG CCTGCAGAAGA CGTGCAGAAG CCAGTCCTT TGGTGTCTAC TTAGACTGGA TGTTCACTCT CCTGGACTAC CTGGACTAC CTGGATTCAA	41	51 ACTGGAAGTC CCCTCCAGGC GGGGAGAGAC CAGCCCTGTT TCCTGACAGC TCCAGAAGGA CATCGCCAGT CTGCTGGCAC TGCCTCACAC TCTCTCCCGG ACTCCTCGGGG GCAAATATGT ACACCCTAAGC AACCCTAAGC GTCACTTAAT GAAATATTGT	60 120 180 240 300 420 480 540 660 720 840 900 960 1020 1080
60 65	1 AGGAATCTGC ATCGGGCAGA CATGAGGATCATC CGAGAAGACG GGAGGCCTGCT CAGCCTCCCC CAGCTGCCTCC CTCGATCACC CAACATCACA CGAGAGATCACA GGGTGACTCC CCAGGATCACA GGGTGACTCC ACCCTCCATT ACCCTCCATT ACCCTCCATT ACCCTCCATT CAAGACCCTC AATCAACCTG GACTCTGGA	26-874 11 GCTCGGGTTC GGTCTCACAG CTGCAGTTCA AAGGGGTTCG CGGCTACTCT CTCAAGCCC CACAAAAGAC TGGGCTGTCC AACAAAGAC GACACATCA GACACATCA GACACATCA GACACATCA GACACATCA GACACATCA GAGAACATCA CAGGGAACA TCACAACATCA TACGAACATTA ATGACAACACA ATGACAACACA	21 21 CGCAGATGCA CAGCCAAGGC AGTGCAAGCC AGTGCAAGCC GTGGGCGAC CCTACATAGT GGACAGCCAC ACCGCAATGA ACCGCCAATGA CACCCTTACA ACTGTTGCACA CCGACACTCAC CCATCATTGA TGTGTGCCAG TGGTCTGTAA GTTTGGGCCA CTTTGGGCCA CTTTGGGCCA CTTTGGGCCC TCATGAGACCA CTTTTGGGCCC CTTTGGGCCC CTGGTGGACC CTGGTTGTTT	31 GAGGTTGAGG ACCTGGGAACA TCACTCCCAG GCTCATCGCC TCACCTGGGG TCACCTGCGG TCACCTGCTC CCATCATCTC CCTCACCCC CCACCAGAG CCTCACCCC CCACCAGAG CCAGCAGGAA CCAGTCTTT TGGTGTCTAC TTAGACTGGT TGTTCACTCT CCTGGACTAC	41 TGGCTGCGGG CGCTCCTCCC GGGCTTGTAG CCCTGGCAGC CCACCCCG GTGAAGATGC CGCTGTGTAC CAGTTACGCC TGTGAGAAC GGGGGCAAGG CAAGGCATTA ACGAAAGTCT CCCACCCACC GTTAATAAGA AGGAGATTCT ACGAGAGCTT ATCCCCAGCCT TCCCCAGCCC TCCCAGCC	51 ACTGGAAGTC CCCTCCAGGC GGGGAGAGAC CAGCCCTGTT TCCTGACAGC TCCAGAAGGA CATCGCCAGT CTGCTGGCAC TGCCTCACAC TCTCTCCCGG ACTCCTCGGGG GCAAATATGT ACACCCTAAGC AACCCTAAGC GTCACTTAAT GAAATATTGT	60 120 180 240 300 360 420 480 540 600 720 780 840 900 900 1020

SEQ ID NO:34 PBA6 PROTEIN SEQUENCE Protein Accession #: NP_006844

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	MRTIOLILLA	LATCINGGET	 RIIKGFECKP	HSOPWOAAT.R	FETTILCAT	I.TADRWI.I.TA	60
5			EGCEOTRTAT				120
	SITWAVRPLT	LSSRCVTAGT	SCLISGWGST	SSPQLRLPHT	LRCANITIE	HQKCENAYPG	180
	NITDTMVCAS DWIOETMKNN		GDSGGPLVCN	QSLQGIISWG	QDPCAITRKP	GVYTKVCKYV	240
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25			CACTTCCAAA				600
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30 .	ACCATAAAAG	AGCTGGAATC	GATTATAAGC	AAAAGGAATA	TTCAATTTTC	CTGCAAGAAT	900
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55	1 ATGTCCTTC	: 1-3315 11 GGGCAGCCAG	(underlined seque 21 GCTCAGCATG	31 AGGAACAGAA	41 GGAATGACAC	51 TCTGGACAGC	60
55	1 ATGTCCTTTC ACCCGGACCC	: 1-3315 11 GGGCAGCCAG TGTACTCCAG	(underlined seque 21 GCTCAGCATG CGCGTCTCGG	31 AGGAACAGAA AGCACAGACT	41 GGAATGACAC TGTCTTACAG	51 TCTGGACAGC TGAAAGCGAC	120
55	1 ATGTCCTTTC ACCCGGACCC TTGGTGAATT	11 1-3315 GGGCAGCCAG TGTACTCCAG TTATTCAAGC	(underlined seque 21 GCTCAGCATG CGCGTCTCGG AAATTTTAAG	31 AGGAACAGAA AGCACAGACT AAACGAGAAT	41 GGAATGACAC TGTCTTACAG GTGTCTTCTT	51 TCTGGACAGC TGAAAGCGAC TACCAAAGAT	120 180
	1 ATGTCCTTTC ACCCGGACCC TTGGTGAATT TCCAAGGCCA ACCCAGATCA	1.3315 11	(underlined seque 21 GCTCAGCATG CGCGTCTCGG AAATTTTAAG GTGCAAGTGT GAAATGGAAC	31 AGGAACAGAA AGCACAGACT AAACGAGAAT GGCTATGCCC TACAAGAAAC	41 GGAATGACAC TGTCTTACAG GTGTCTTCTT AGAGCCAGCA ACACCAAGGA	51 TCTGGACAGC TGAAAGCGAC TACCAAAGAT CATGGAAGGC ATTTCCTACC	120
55· 60	1 ATGTCCTTTC ACCCGGACCC TTGGTGAATT TCCAAGGCCA ACCCAGATCA GACGCCTTTG	1.3315 11 GGGCAGCCAG TGTACTCCAG TGTATTCAAGC CGGAGAATGT ACCAAAGTGA GGGATATTCA	(underlined seque 21 GCTCAGCATG CGCGTCTCGG AAATTTTAAG GTGCAAGTGT GAAATGGAAC GTTTGAGACA	31 AGGAACAGAA AGCACAGACT AAACGAGAAT GGCTATGCCC TACAAGAAAC CTGGGGAAGA	41 GGAATGACAC TGTCTTACAG GTGTCTTCTT AGAGCCAGCA ACACCAAGGA AAGGGAAGTA	51 TCTGGACAGC TGAAAGCGAC TACCAAAGAT CATGGAAGGC ATTTCCTACC TATACGTCTG	120 180 240 300 360
	1 ATGTCCTTTC ACCCGGACCC TTGGTGAATT TCCAAGGCCA ACCCAGATCA GACGCCTTTG TCCTGCGACA	1.3315 11	(underlined seque 21 GCTCAGCATG CGCGTCTCGG AAATTTTAAG GTGCAAGTGT GAAATGGAAC GTTTGAGACA AATCCTTTAC	31 - AGGAACAGAAT AGCACAGACT AAACGAGAAT GGCTATGCCC TACAAGAAAC CTGGGGAAGA GAGCTGCTGA	41 GGAATGACAC TGTCTTACAG GTGTCTTCTT AGAGCCAGGA ACACCAAGGA AAGGGAAGTA CCCAGCACTG	51 TCTGGACAGC TGAAAGCGAC TACCAAAGAT CATGGAAGGC ATTTCCTACC TATACGTCTG GCACCTGARA	120 180 240 300 360 420
	1 ATGTCCTTTC ACCCGGACCC TTGGTGAATT TCCAAGGCCA ACCCAGATCA GACGCCTTTG TCCTGCGACA ACACCCAACC	1.3315 11	(underlined seque 21 GCTCAGCATG CGCGTCTCGG AAATTTTAAG GTGCAAGTGT GAAATGGAAC GTTTGAGACA	31 AGGAACAGAA AGCACAGACT AAACGAGAAT GGCTATGCCC TACAAGAAAC CACGGCAAGA GAGCTGCTGA GGCGCCAAGA	41 GGAATGACAC TGTCTTACAG GTGTCTTCTT AGAGCCAGGA ACACCAAGGA AAGGGAAGTA CCCAGGACTG ACTTCGCCCT	51 TCTGGACAGC TGAAAGCGAC TACCAAAGAT CATGGAAGGC ATTTCCTACC TATACGTCTG GCACCTGARA GAAGCCGCGC	120 180 240 300 360
60	1 ATGTCCTTTC ACCCGGACCC TTGGTGAATT TCCAAGGCCA ACCCAACC TCCTGCGACA ACACCCAACC ATGCGCAAGA ACGCGAGGA	1.3315 11 GGGCAGCCAG TGTACTCCAG TGTACTCCAG TGTATTCAAGC CGGAGAATGTA ACCAAAGTGA GGGATATTCA CGGACGCGGA TGGTCATTTC TCTTCAGCCG CCCATTATGG	(underlined seque 21 GCTCAGCATG GCGGTCTCGG AAATTTTAAG GTGCAAGTGT GAAATGGAAC GTTTGAGACA AATCCTTTAC TGTGACCGGG GCTCATCTAC CCTGATGAAG CCTGATGAAG	31 AGGAACAGAA AGCACAGACT AAACGAGAAT GGCTATGCCC TACAAGAAAC CTGGGGAAGA GAGCTGCTGA GGCGCCAAGA ATCGCGCAGT TACATCGGGG	41 GGAATGACAC TGTCTTACAG GTGTCTTCTT ACACCAAGGA ACACCAAGGA AAGGGAAGTA CCCAGCACTG ACTTCGCCCT CCAAAGGTGC AGGTGGTGAG	51 TCTGGACAGC TGAAAGCGAC TACCAAAGAT CATGGAAGGC ATTTCCTACC TATACGTCTG GCACCTGARA GAAGCCCCGC TTGGATTCTC AGATAACACC	120 180 240 300 360 420 480 540
	1 ATGTCCTTTC ACCCGGACCC TTGGTGAATT TCCAAGGCCA ACCCGAATCA GACGCCTTTG TCCTCCGACA ACACCCAACC ATCGCCAAGC ATCGCGAGGCA	1-3315 11 GGGCAGCCAG TGTACTCCAG TGTACTCAGC CGGAGAATGT ACCAAAGTGA ACCAAAGTGA CGGACGCGGA TGGTCATTTC TCTTCAGCCG CCCATTATGG GTTCAGAGGA	21	31 	41 GGAATGACAC TGTCTTACAG GTGTCTTCTT AGAGGCAGGA ACACCAAGGA AAGGGAAGTTA CCCAGCACTG ACTTCGCCCT CCAAAGGTGC AGGTGGTGAG TAGCAGCTTG	51 TCTGGACAGC TGAAAGCAC TACCAAAGAT CATGGAAGCC TATACCTCTG GCACCTGAAA GAAGCCGCGC TTGGATTCTC AGATAACACC GGGCATGGTC	120 180 240 300 360 420 480 540 600 660
60	1 I ATGTCCTTTC ACCGGACCC TTGGTGAATT TCCAAGGCCA ACCCAGATCA GACGCCTTTG GACACCAACC ATGCGCAAGA ACACCCAACC ATGCGGAGGCA ATCAGCAGGA TCCAACCGG	1.3315 11	21 GCTCAGCATG CCCGTCTCGG AAATTTTAAG GTGCAAGTGT GAAATGGAAC GTTTGAGACA AATCCTTTAC TGTGACCAGG GCTCATCTAC CCTGATGAAG CATATTTTAC GAATATTGTG CAGGAATTGC CAGGAATTGC	31 AGGAACAGAA AGCACAGACT AAACGAGAAA GGCTATGCCC TACAAGAAAC CTGGGGAAGA GAGCTGCTGA GGCGCCAAGA ATCGCGCAGT TACATCGGGG GCCATTGGCA GATGCTGAAG	41 GGAATGACAC TGTCTTCTTA AGAGCCAGCA ACACCAAGGA ACACCAGCACTG ACTTCGCCCT CCAAAGGTGC AGGTGGTGAG AGGTGGTGAG GCTATTTTTT	51 TCTGGACAGC TGAAAGCCAC TACCAAAGAT CATGGAAGGC ATTTCCTACC TATACCTCTG GCACCTGAAA GAAGCCGCGC TTGGATTCTC AGATAACACC GGGCATGGTC AGCCAGTAC	120 180 240 300 360 420 480 540 600 660 720
60	1 ATGTCCTTTC ACCCGGACCC TTGGTGAATT TCCAAGGCCA ACCCAGATCA GACGCCTTTG TCCTGCGACA ACACCCAACC ATGCGCAAGA ACGCGAGGCA ATCAGCAGGGA TCCAACCGGG CTTATGGATG CTGCTCGTGG	1.3315 11 GGGCAGCCAG TOTACTCCAG TOTACTCCAG TOTACTCCAG TOTACTCAGCGGA ACGCAAAGTGA TCGTCATTCA CGGACGCGGA TCGTCATTTC TCTTCAGCCG CCCATTATGG GTCAGCAGGA ACACCCTCAT ACTTCACAAG ACAATGGCTG	(underlined seque 21 GCTCAGCATG CCCGTCTCGG AAATTTTAAG GTGCAAGTGT GAAATGGAAC GTTTGAGACA AATCCTTTAC TGTGACCGGG GCTCATCTAC CCTGATGAAG GAATATTGTG CAGGAATTGC AGATCCACTG TCATGGACAT TCATGGACAT	31 AGGAACAGAA AGCACAGACT AAACGAGAAT GGCTATGCCC TACAAGAAAC CTGGGGAAGA ATCGCGCAGT TACATCGGG GCCATTGGCA GATGCTGAGG TATATCCTGG CCCACTGTCG	41 GGAATGACAC TGTCTTACAG GTGTCTTCTT AGAGCCAGGA ACACCAAGGA ACACGACTA CCCAAAGGTGC CCAAAGGTGC AGGTGGTGAG TAGCAGCTTG GCTATTTTTT ACAACAACCA AAGCAAACCT	51 TCTGGACAGC TGAAAGCAC TACCAAAGAT CATGGAAGGC ATTTCCTACC TATACGTCTG GCACCTGARA GAAGCCGCGC TTGGATTCTC AGATACACC GGGCATGGTC AGCCCAGTAC AGCCCAGTAC AGCCCAGTTC CCACACATTTG CCGGAATCAG	120 180 240 300 360 420 480 540 660 720 780 840
60 65	1 ATGTCCTTTC ACCCGGACCC TTGGTGAATT TCCAAGGCCA ACACCCAACC ATGCGAACA ATGCGCAACA ATGCGCAAGA ATCAGCAGGA TCCAACCGG CTTATGGATG CTGCTCGTGG CTTATGGATG CTGCTCGTGG CTTAGGAGA	1.3315 11	(underlined seque 21 GCTCAGCATG GCGCTCTCG AAATTTTAAG GTGCAAGTGT GAAATGGAAC GTTTGAGCAC GTTTGAGCAC GCTCATCTAC CCTGATCAAG GCAATATTGTG CAGGAATTGTG CAGGAATTGC AGATCACTG GCACTATTA GCGCACTATT	31 	41	51 TCTGGACAGC TGAAAGCAC TACCAAAGAT CATGGAAGCC TATACCTCTG GCACCTGAAA GAAGCCCCGC TTGGATTCTC AGGTAACACC GGGCATGGTC AGCCCACTAC CACACATTTG CCGGAATCAG CAAGATCCCC	120 180 240 300 360 420 480 540 660 720 780 840 900
60	1 I ATGTCCTTTC ACCCGGACCC TTGGTGAATTA ACCCAGATCA ACCCCAACC ACGCCAACC ATGCGCAAGA ACGCGAGGA ATCAGCAGGA TCCAACCGGG CTTATGGATG CTGCTCGTGG CTAGGAGAGT ATTGTCTGTTT	1.3315 11 GGGCAGCCAG TTATTCCAG TTATTCAAGC CGGAGAATGT ACCAAAGTGA TGGTCATTTC TCTTCAGCCG CCCATTATGG GTTCAGAGGA ACACCTCAT ACTTCACAAG ACAATGCTGA ACAATGCTGA TTGCCCAAGG TTGCCCAAGG	21 GCTCAGCATG CCCGTCTCGG AAATTTTAAG GTGCAAGTGT GAAATGGAAC GTTTGAGACA CTTTGACCGG GCTCATCTAC CCTGATGAAC CCTGATGAAC CAAATTTTAC CAGAAATTGTG CAGGAATTGTG CAGGAATTGTG CAGGAATTGTA AGGTGGAAAA AGCTGGACAAT	31 AGGAACAGAA AGCACAGACT AAACGAGAAAC CTGGGGAAGA GAGCTGCTGA GCGCCCAAGA ATCGCGCAGT TACATCGGG GCCATTGGCA GATGCTGAGG TATATCCTGC CCACTGTCG CAAGATTCCA GAGACTTTGA	41 GGAATGACAC TOTOTTACAG GTGTOTTOTTA ACACCAAGGA AAGGGAAGTA ACTCGCCCT CCAAAGGTGC AGGTGGTGAG GCTATTTTTT ACAACAACCA AAGCAAACCA AAGCAAACCA AAGCAAACCA AAGCAAACCA AAGCAAACCA	51 TCTGGACAGC TGAAAGCAC TACCAAAGAT CATGGAAGCC TATACCTCTG GCACCTGAAA GAAGCCCCGC TTGGATTCTC AGGTAACACC GGGCATGGTC AGCCCACTAC CACACATTTG CCGGAATCAG CAAGATCCCC	120 180 240 300 360 420 480 540 600 720 780 840 900 960
60 65	1 ATGTCCTTTC ACCCGGACCC TTGGTGAATT TCCAAGGCCA ACACCCAACC ATGCGCAACA ACGCCAACA ACGCCAACC ATCAGCAACC ATCAGCAGA ACCAACCGACC ATCAGCAGGA ACCAACCGGGCAGCA ATCAGCAGGA ATCAGCAGGA TCCAACCGGG CTTATGGATG CTGCTCGTGG CTAGTGGTTGTTTTGTGTGTTTT AAAATAAAA AGCCTGGTGG	1.3315 11 GGGCAGCCAG TOTACTCCAG TOTACTCCAG TOTACTCAGC GGACAGCGGA ACGCATATTCA CCGAGCAGCGGA ACACCCTCAT ACTICAGCAG ACACCCTCAT ACTICACAAG ACACCTCAT ACTICACAAG ACACCTCAT ACTICACAAG ACACCTCTCT ATACCCAGA ACACCTCTCTACAAG ACACTCTCTACAAG ACACTCTCTACAAG ACACTCTCTACAAG ACACTCTTCTGTT AGGTGGAGGA	(underlined seque 21 GCTCAGCATG GCGTCTCGG ANATTTTAG GTGCAAGTGT GAAATGGAAC GTTTGACCGG GCTCATCTTAC CCTGATGAAG GAATATTGTG CAGGAATTGC AGATCCACTG TCATGGACAT GCGCACTATT AGGTGGAAAA AGGTGGTGGAAA AGGTGGTGGAAA TGCCCTGACACA	31 AGGAACAGAA AGCACAGACT AAACGAGAAT GGCTATGCCC TACAAGAAAC CTGGGGAAGA ATCGCGCAGT TACATCGGGG GCCATTGGCA GATGCTGAGG TATATCCTGG CCCACTGTCG CAAGACTTTCA GAGACTTTCA GGCTCGGGCC TCTTCTGCCG	41	51 TCTGGACAGC TGAAAGCAA TACCAAAGAT CATGGAAGGC ATTTCCTACC TATACGTCTG GCACCTGARA GAAGCCCCGC TTGGATTCTC AGATAACACC GGGCATGGTC AGCCCAGTAC CACACATTTG CCGGATCAG CAAGATCCC TACCTCCATC TGTGATCGCT GCTGGTCCCC	120 180 240 300 360 420 480 540 660 720 780 840 900 960 1020 1080
60 65	1 ATGTCCTTTC ACCGGACCC TTGGTGAATT TCCAAGGCCA ACCCAGATCA ACACCCAACC ATGCGCAACC ATGCGCAAGC ATGCGCAGGA ACAGCAGGA TCCAACCGG TCCAACCGG CTTATGGATG CTGCTCGTGG CTGAGGAAGT ATTGTGTGTT AAAAATAAAA AGCCTGGTGG TTTTTACCCC	1.3315 11 GGGCAGCCAG TITACTCCAG TITACTCAAG CGGAGAATGT ACCAAAGTGA ACGACGCGGA TGGTCATTTC TCTTCAGCCG CCCATTATGG GCTTCAGAGGA ACACCCTCAT ACTTCACAGG ACACCCTCAT ACTTCACAGG TTCCTCAGGCT TTCCTTGTGT TTCCTTGTGT TTCCTTGTGT TGCCTAGGGA ACACCCTCAT ACTTCACAGGCA ACACCCTCAT ACTTCACAGGCA ACACCCTCAT ACTTCACAGGCA ACACCCTCAG ACACCCTCAGGCA ACACCCTCAGAGGA ACACCCTCAGAGGA ACACCCTCAGAGGA ACACCCTCAGAGGA ACACCCCTCAGAGGA ACACCCCTCAGAGGA ACACCCCTCAGAGGA ACACCCCTCAGAGGA ACACCCCTCAGAGGA ACACCCCTCAGAGGA	(underlined seque 21 GCTCAGCATG CCGGTCTCGG AAATTTTAAG GTGCAAGTGT GAAATGGAAC GTTTGAGCAC GTTTGAGCAC CCTGATCAC CCTGATCAC CCTGATCAC CAGGAATTGT CAGGAATTGT CAGGACAT CCGCACTATT AGGTGGACAT AGGTGGACAA GGTGGTGGACA CCCGGCTGCCT CCGGCTGCCC	31 AGGAACAGAA AGCACAGACT AAACGAGAAC GGCTATGCCC TACAAGAAAC CTGGGGAAGA ATCGCGCAAGA ATCGCGCAGT TACATCGGG GCCATTGGCA GATGCTGAGG CCACTGTCG CCACTGTCG CAAGATTCCA GAGACTTTGA GGCTCGGGCC TCTTCTGCCG GAGGAGGAGA	41 GGAATGACAC TGTCTTCTT AGAGCCAGCA ACACCAAGGA ACACCAAGGA ACACCATC CCAAAGGTGC ACGTGCTTG CCTATTTTTT ACAACAACCA ACGACAC ACGACACTG ACTTCGCCT GCTATTTTTT ACAACAACCA ACGACACAC ACTTCGCCA ACGACACC ACTTCGCCCT ACTACTTCTT ACAACAACCA ACGACAAC ACATCGCTGG AAGCCATCAA ACGATCGCTGA ACGATCGCTGA ACGATCGCTGA ACGATCGCTGA ACGATCGCTGA ACGAGGAGAA CTGAGGAGAA	51 TCTGGACAGC TGAAAGCCAC TACCAAAGAT CATGGAAGGC ATTTCCTACC TATACCTCTG GCACCTGAAA GAAGCCGCGC TTGGATTCC AGATAACACC AGGCATGGTC AGCCAGTAC CACACATTTG CCAGCATCAG CAAGATCCC TACCTCCATC TGTGATCGCT GCTGGTGGCG GATCAAATGG	120 180 240 300 360 420 480 540 600 720 780 840 900 900 1020 1080 1140
60 65 70	1 ATGTCCTTTC ACCGGACCC TTGGTGAATT TCCAAGGCTA ACCCAGATCA ACACCAACC ATGCGCAAGA ACAGCCAACC ATGCGCAGGA TCCAACCGGG CTTATGGATG CTGCTCGGGC CTAGAGAGT ATTGTGTTT AAAATAAAA AGCCTGGTGG TTTTTACCCC CTCAAAGAAA	1.3315 11	(underlined seque 21 GCTCAGCATG CCCGTCTCGG AAATTTTAAG GTGCAAGTGT GAAATGGAAC GTTTGAGACA CCTGATGAAC CCTGATGAC GAAATTTTAC TGTGACCGG GCTCATCTAC CCTGATGACA TCATGACAT AGATCACTG TCATGGACAT AGGTGGAAAA GGTGGTGGAAA GGTGGTGGAAA GCTGGTGGAAA GCTGGTGGAAA CCGGCTGCCT TTCTCACCTA	31 AGGAACAGAA AGCACAGACT AAACGAGAAC GGCTATGCCC TACAAGAAAC CTGGGGAAGA ATCGCGCAGT TACATCGGG GCCATTGGCA GATCCTGAGG TATATCCTGC CCACTGTCG CAAGATTCCA GAGACTTTTCA GGCTCGGGCC TCTTCTGCCG GAGGAGAGA TTAACAGTTA	41 GGAATGACAC TOTOTTACAG GTGTCTTCTTA AGAGCCAGCAC ACACCAAGGA AAGGGAAGTA ACTCGCCCT CCAAAGGTGC AGGTGGTGAG GCTATTTTTT ACAACAACCA AAGCAAACCA AAGCAAACCA AAGCAAACCA AAGCAAACCA AAGCAAACCA CTAAGGAGAT CCAAGGAGAT TCAAAGGAGTT TTAAAATGGA	51 TCTGGACAGC TGAAAGCGAC TACCAAAGAT CATGGAAGCC ATTTCCTACC TATACGTCTG GCACCTGARA GAAGCCGCGC TTGGATTCTC AGGATACACC AGGCCAGTAC CACACATTTG CCGGAATCAG CAAGATCACC TACCTCCATC TGTGATCGCT GCTGGTCCGC GATCAAATGG AGAAGCTCGG	120 180 240 300 360 420 480 540 660 720 780 900 960 1020 1080 1140 1200
60 65	1 ATGTCCTTTC ACCCGGACCC TTGGTGAATT TCCAAGGCCA ACACCCAACC ATGCGACAAC ACGCCAACC ATCGCCAACA ATCAGCAGAC ATCAGCAGGA TCCAACCGGACGA TCCAACCGGACC ATCAGCAGGA ATCAGCAGGA TCTATGGATG CTTATGGTTGT AAAATAAAA AGCCTGGTGG TTTTTACCCC CTCAAAGAAAC CAAGAAATAG CAAGAAAAGG	1.3315 11	underlined seque 21 GCTCAGCATG CCGCTCTCGG ANATTITAAG GTGCAAGTGT GAAATGGAAC GTTTGAGCAC TGTGACCAGG GCTCATCTAC CCTGATGAG CAGAATTGTG CAGGAATTGTG CAGGAATTGTG CAGGAATTGTG CAGGAATTGTG CAGGACAT CCGCACTATT AGGTGGAAA GCTGGTGGAA TGCCCTGACA CCGGCTGCCT TTCTCACCTA CATCTCCTAC TGGGCAGCTTG	31 AGGAACAGAA AGCACAGACT AAACGAGAAC CTGGGGAAGA GGCTGCTGA ATCGGGCAAGA ATCGCGCAGT TACATCGGG GCCATTGGCA GATCCTGAG GATCCTGAG GCACTGTCG CAAGATTCCA GAGACTTCCA GAGACTTCCA GAGACTTCCA GAGACTTCCA GAGACTTCCA GAGACTTCAC GAGAGAGA TTAAACAGTA CCTCTAACA AAGCTTCTGC	41 GGAATGACAC TGTCTTACAG GTGTCTTCTT AGAGCCAGCA ACACCAAGGA AAGGGAAGTA CCCAGCACTG CCTAAAGGTGC AGGTGGTGAG TAGCAGCTTG ACATCACCA AAGCAAACCA AAGCAAACCA AAGCAACCA AAGCAACCA TCAAGGAGAA CTGAGGAGAA CTGAGGAGTTG TTAAAATTGG TGAGAGTTGA AAGCCTTCAG TGGAGTTGAA	51 TCTGGACAGC TGAAAGCAT CATGGAAGGC ATTTCCTACC TATACGTCTG GCACCTGARA GAAGCCGCGC TTGGATTCTC AGATACACC GGGCATGGTC AGCCAGTTC CCACACATTTG CCAGATCAG CAAGATCCC TACCTCCATC GCTGGTCCGC GATCAAATGG CACACATTGG CCAGATCAG CCAGCAGC CACCAGTCAG CCACCATGAC CCACCATGAG CCACCTGAG CCACCTGAG CCACCTGAG CCACCTGAG	120 180 240 360 420 480 540 660 720 840 960 960 1020 1140 1200 1260 1320
60 65 70	1 ATGTCCTTTC ACCGGACCC TTGGTGAATT TCCAAGGCCA ACCCAGATCA ACACCAACC ATGCGCACA ACACCCAACC ATGCGCACA ACACCCAGC TCCAACCGG TCCAACCGG CTTATGGATG CTGCTCGTGG CTGAGGAAGT AAAATAAAA ACCCTGCTGC TTTTTACCC CTCAAAGAATA GATGAAATA CAAGAAATA CAAGAAATA CAAGAAATA CAAGAAATA CAAGAAATG TTAGCCAATG	1.3315 11 GGGCAGCCAG TITACTCCAG TITACTCAAGC CGGAGATATTC ACCAAAGTGA ACGAACGCGA TGGTCATTTC TCTTCAGCCG CCCATTATGG GCTTCAGAGGA ACACCCTCAT ACTTCACAAG ACATCCTGA TTCCCTAGTG TTCCCTAGTG ACATCCTGA TTCCTTGTGT TTCTCGAATG TCACAGCACC TCCTCGATTC TCACAGCACC TTCTCGATTG TGACAGCACTC TCACAGCATTC TCACAATG TCACAGCATTC TCACAATG TCACAGCATTC TCACAATG TCACAGCATTC TCACAATG TCACAGCATTC TCACAATG TCACAGCATTC TCACAATG TCACACTC TACACTGGAA ATGACATTTT	(underlined seque 21 GCTCAGCATG GCGCTCTGG AAATTTTAAG GTGCAAGTGT GAAATGGAAC GTTTGAGCAG GCTCATCTAC CCTGATCAAG GCTCATCTAC CAGGAATTGT CAGGAATTGT AGGTGACAT GCGCACTATT AGGTGGAAAA GGTGGTGGAAA GGTGGTGGAAA GCCCTGACC CCGCTGCCT TTCTCACCTA CAGCACTGCCT CAGCACTATC CATCTCCTAC CATCTCCTAC CACCACTATC CACCACTATC CACCACTATC CACCACTACC CACCACTACAC CCCCCACTACC CCCCCACTACC CACCACTACC CACCACTACC CACCACTACC CACCACTACC CCCCCACTACC CACCACTACC CACCACTACC CACCACTACC CACCACTACC CCCCCACTACC CCCCCACTACC CACCACTACC CCCCCACTACC CCCCCACTACC CCCCACTACC CCCCCACTACC CCCCCACTACC CCCCCACTACC CCCCCCCC	31 AGGAACAGAA AGCACAGACT AAACGAGAAC GGCTATGCCC TACAAGAAAC CTGGGGAAGA ATCGCGCACT TACATCGGG GCCATAGCC GATGCTGAG GATGCTGAG GATGCTGAG CCACTGTCG CAAGATTCCA GAGACTTTGA GGCTCGGGCC TTTCTCCCC GAGGAGGAGA TTAACAGTTA CCTCTATACA CCTCTATACA CCCCCTATCTCCCC CGCGATGGC CCCCCTCTCTCCCC CAGGAGGAGA CTAACAGTTA CCTCTTATACA AAGCTTCTCCC CGCCGATGGG CCCCCATGGGC CCCCGATGGG	41 GGAATGACAC TGTCTTCTACAG GTGTCTTCTTACAG AGAGCCAGCAC ACACCAAGGA AAGGGAAGTA ACTCGCCCT CCAAAGGTGCA AGGTGGTGAG GCTATTTTTT ACAACAACCAA AAGCAAAGCTA AAGCAAAGCTA AAGCAAGCTAC AAGCAAGCTGA AAGCAAGCTTGA AGATCGCTGA AGATCGCTGA ACTATGGTGG AAGCCTTCAA AGATCGCTGA AAGCCTTCAA AGGTGGAA AGGTCTGCAA AGGTCGCAA AGGTCGCAA AGGTCGCAA	51 TCTGGACAGC TGAAAGCCAC TACCAAAGAT CATGGAAGGC ATTTCCTACC TATACOTCTG GCACCTGAAA GAAGCCCGC TTGGATTCTC AGGATACACC CAGCATTAC CACACATTG CCGGAATCAG CAAGATCCC TACCTCCATC TGTGATCGCT GCTGGTGCGC GATCAAATGG AGAAGCTGGG CACCACTGAG CACCACTGAG CACCACTGAG	120 180 240 360 420 480 540 660 720 960 1020 1140 1260 1320 1380
60 65 70	1 ATGTCCTTTC ACCCGGACCC TTGGTGAATT TCCAAGGCCA ACCCAGATCA ACCCCAACC ATGCGCAAGA ACGGGAGGCA ATCAGCAGGGA TCCAACCGGG CTTATGGATG CTGCTGTGG CTAGCAGGT TATGTGTTT AAAATAAA AGCCTGGTGT TTTTACCCC CTCAAAGAAA GATGAAATC CAAGACAAG TTAGCAAG TTAGCAAG TTAGCACT TTAGCAC TTAGCAAGT TTTTACCCC TCCAAAGAAA GATGAAATT CAAGACAAGG TTAGCCAATG TTAGCCAATG TTAGCCAATG	1-3315 11 GGGCAGCCAG TTATTCCAG TTATTCAAGC CGGAGAATGT ACCAAAGTGA TGGTCATTTC TCTTCAGCCG CCCATTATGG GTTCAGAGGA ACACCTCAT ACTTCACAAG ACAATGCTGA TTGCCCAAG TTCCTTGTGT AGGTGGAGGA TTCCTTGTGT TGGCCAATGC TTCTCGAATG TTCTCGAATG TTCTCGAATG TGAGCAATGCT TCTCGAATG TGAGCAATGCT ATACCTCAATG TGAGCAATGC ATACTGTAACTGGAATG TGAGCAATGC ATAACTGCAATG TGAGCAATGC ATAACTGCAATG TGAGCATTTT CGGCTCTCAT	(underlined seque 21 GCTCAGCATG CCCGTCTCGG AAATTTTAAG GTGCAAGTGT GAAATGGAAC GTTTGAGACA CCTGATGAAC GCTCATCTAC CCTGATGAAG GAAATTTTA GGAAATTGTG CAGGAATTGTG CAGGAATTGTG CAGGAATTGTG CAGGAATTGTG CAGGACATAT AGGTGGAAAA GGTGGTGGAA GGTGGTGGAA GGTGGTGGAA CCGGCTGCCT TTCTCACCTA CATCTCCTAC CACCAATGAC AAAGGACAGA AAAGGACAGA	31 AGGAACAGAA AGCACAGACT AAACGAGAAT GGCTATGCCC TACAAGAAAC CTGGGGAAGA ATCGCGCAGT TACATCGGG GCCATTGGCA GATGCTGAGG CACATTGCC CAAGATTCCA GAGACTTTCA GGCTCGGGCC TTTCTGCCG GAGGAGAGA TTAACAGTTA GCTCTATACA AAGCTTCTGC CCCCGATGGG CCCCAAGTTTG	41	51 TCTGGACAGC TGAAAGCGAC TGAAAGAT TACCAAAGAT CATGGAGCC ATTTCCTACC TATACGTCTG GCACCTGARA GAAGCCGCGC TTGGATTCTC AGATAACACC GGGCATGAT CACACATTTG CCGGAATCAG CAAGATCACC TACCTCCATC TGTGATCGCT GCTGGTCGCG CATCAAATGG AGAAGCTCGG CACCAGTGAG CCCTTCAAGAA TCTGGAGAAT	120 180 240 300 360 420 540 600 660 720 780 840 900 1020 1140 1200 1260 1380 1440
60 65 70	1 ATGTCCTTTC ACCCGGACCC TTGGTGAATT TCCAAGGCCA ACCCAGATCA GACGCCTTTG TCCTGCGACA ACACCCAACC ATCAGCAGAT ACACCCAACC ATCAGCAGGA ACCAACCGG CTTATGGATG CTGACAGGA ATCAGCAGT ATTGTGTGT AAAATAAAA AGCCTGGTGG TTTTTACCCC CTCAAAGAAA GATGAAATTG CAAGCAAGG TTAGCCAATT GAAGCAATTG GTCATGTTTA GGCTTGTAC GTCATGTTTA GGCTTGTAC GTCATGTTTA GGCTTGAACCA GTCATGTTTA GGCTTGAACCA GTCATGTTTA GGCTTGAACCA GTCATGTACCA GTCATGTACCC TTAGCCAATTG GTCATGTACCC TTAGCCAATTG GTCATGTACCC	1-3315 11	(underlined seque 21 GCTCAGCATG CGCGTCTCGG ANATTTTAAG GTGCAAGTGT GAAATGGAAC GTTTGAGACA AATCCTTTAC CCTGATGAAG GAATATTGTG CAGGAATTGC GAAATGTAC CCTGATGAAG GAATATTGTG CAGGAATTGC AGATCACTG TCATGGACAT TCATGGACAT TCCTGACAT TCCTGACAT TCCTGACAT TCCTGACAT TCCTGACAT TCCTCACCTA CATCTCCTAC TGGGCAGCTGCT CATCTCCTAC TGGGCAGCTGC CACCAATTGA AAAGGACAGA TCTCACCCAT	31 AGGAACAGAA AGCACAGACT AAACGAGAAC GGCTATGCCC TACAAGAAAC CTGGGGAAGA ATCGCGCAGT TACATCGGG GCCAATGCCA CACACTGCCA CACACTGCCA CAGACTTCCA CAGACTTCCA CAGACTTCCA CAGACTTCCA CAGACTTCCA CAGACTTCCA CAGACTTCCA CAGACTTCCA CAGCACTCCA CAGCACTCCC CACACTCCC CACACTCCC CACACTCCC CACACTCCC CACACTCCC CACACTCCC CACACTCCC CACACTCCC CACACTTCCC CACACCTCCC CACACCTCC CACACC CACACCTCC CACACCTCC CACACCTC CACACC CACACCTC CAC	41 GGAATGACAC TGTCTTACAG GTGTCTTCTTACAG GTGTCTTCTTACAG ACACCAAGGCA ACACCAAGGAA ACACCACTG CCAAAGGTGC AGGTGGTGAG TAGCAGCTTTTT ACAACAACCA AAGCAAACCT ACTATGGTGGA ACGCATCAA AGGATCGCTGA TCAAGGAGATA TCAAGGAGATA TCAAGGAGATA TCAAGGAGAA AGTCTCTCAG TGGAGTTGAACTTCAC TCGAACTCTT TCGGACTCTT TCCGCCTCTT TCTGAACTCTT TCTGAACTCTT TCTGAACTCTT	51 TCTGGACAGC TGAAAGCCAC TACCAAAGAT CATGGAAGGC ATTTCCTACC TATACOTCTG GCACCTGAAA GAAGCCCGC TTGGATTCTC AGGATACACC CAGCATTAC CACACATTG CCGGAATCAG CAAGATCCC TACCTCCATC TGTGATCGCT GCTGGTGCGC GATCAAATGG AGAAGCTGGG CACCACTGAG CACCACTGAG CACCACTGAG	120 180 240 300 360 420 540 600 660 720 780 840 900 1020 1080 1140 1260 1320 1380 1440 1500

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5			CATCAATGCT				1860
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			CTGTGAAGCT				1980
	GTGGAGGCCA	CAGACCAGCA	TTTCATCGCC	CAGCCTGGGG	TCCAGAATTT	TCTTTCTAAG	2040
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3420

3480

3540

3600

3660

3720

3780

3840

3900

3960

SEQ ID NO:42 PBH5 PROTEIN SEQUENCE Prolein Accession #: NP 005836

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45	TREAVVARLA LTELKLAIDG WVFNGRPHCF EGYVYVGLYL VRTDLNYIAA Nucleic Acid Acce Coding sequence: 1 GGAGCAGCCCT	DDMLEKLPPD TIIMSENLQD WMTGFFNPQG EGAGWDERNM VDLRTAQTPE sslon #: AB033 68-3345 11	YVPFEVKERL ALICMFDARI FLTAMRQEIT KLIESKPKVL E HWVLRGVA 036 (underlined sequ 21 	OKMGPFOPMN PAWWKKASWV RANKGWALDN FELMPVIRTY LL CDVK SEC ences correspond 31 CACTACCCCT	IFLRQEIDRM FSTLGFWFTE MVLCNEVTKW AENNTLRDPR 2 ID NO:47 PDG5 to start and stop of 41 1 CAGGGGTTGC	QRVLSLVRST LIERNSQFTS MKDDISTPPT FYSCPIYKKP DNA SEQUENCE odons) 51 TTTCAGATAA	180 240 300 360
45 50	TREAVVARLA LTELKLAIDG WVFNGRPHCF EGYVYVGLYL VRTDLNYIAA Nucleic Acid Acce Coding sequence: 1 GGAGCAGCCT AGATGACATG	DDMLEKLPPD TIIMSENLQD WMTGFFNPQG EGAGWDKRNM VDLRTAQTPE SSlon #: AB033 68-3349 11	YVPFEVKERL ALDCMFDARI FLTAMRQEIT KLIESKPKVL E HWVLRGVA 036 0 (underlined sequ 21 ARCCAGAAAAC ATGCTGGCAI	OKMGPFOPMN PAWWKKASWU RANKGWALDN FELMPVIRIY LL CDVK SEC ences correspond 31	IFLROEIDRM FSTLGFWFTE MVLCNEVTKW AENNTLRDPR 2 ID NO:47 PDG5 to start and stop of 41	QRVLSLVRST LIERNSQFTS MKDDISTPPT FYSCPIYKKP DNA SEQUENCE odons) 51 TTTCAGATAA CATCAGCAGC	180 240 300 360 60 120
45	TREAVVARLA LITELKLAIDG WVFNGRPHCF EGVYVYGLYL VRTDLNYIAA Nucleic Acid Acce Coding sequence: 1 GGAGCAGCCT AGATGACATG ACAGCCCATA	DDMLEKLPPD TIIMSENLQD WMTGFFNPQG EGAGWDERNM VDLRTAQTPE sslon#: AB033 68-3349 11 ACAACTTCAC GGAAGGAGAA CCTGAAAACA	YVPFEVKERL ALDCMFDARI FLTAMRQEIT KLIESKPKVL E HWVLRGVA 036 0 (underlined sequ 21 AACCAGAAAC ATGCTGGCAT	OKMGPFOPMN PAWWKKASWV RANKGWALDN FELMPVIRIY LL CDVK SEC ences correspond 31 CACTACCCCT AGATTTCGGA CATGGTTAGT	IFLROEIDRM FSTLGFWFTE MVLCNEVTKW AENNTLRDPR DID NO:47 PDG5 to start and stop of 41 CAGGGGTTGC TCCAGAAAAG GATCCACAAC	QRVLSLVRST LIERNSQFTS MKDDISTPPT FYSCFIYKKP DNA SEQUENCE odons) 51 TTTCAGATAA CATCAGCAGC CATACCATGA	180 240 300 360 60 120 180
45 50	TREAVVARLA LTELKLAIDG WVFNGRPHCF EGYVYYGLYL VRTDLNYIAA Nucleic Acid Acce Coding sequence: 1 GGAGCAGCCT AGATGACATG ACAGCCATA AGATGCAGCT	DDMLEKLPPD TIIMSENLQD WMTGPFNPQG EGAGWDKRMM VDLRTAQTPF sslon #: AB033 68-3345 11 ACAACTTCAC GGAAGGAGAA TCTGGAGCTG TCTGAACACTTCACC TCTGAACCACTTCACC TCTGAACCACTTCACC TCTGAACCACTTCACC TCTGAACCACTTCACC TCTGAACCACTTCACC	YVPFEVKERL ALDCMFDARI FLTAMRQEIT KLIESKPKVL E HWVLRGVA 036 Qunderlined sequence 21 AACCAGAAAC ATGCTGGCA TGGACAATTC AGAAGACAGA	QKMGPFQPMN PAWWKKASWV RANKGWALDN FELMPVIRTY LL CDVK SEC ences correspond 31 CACTACCCCT AGATTTCGCT AGATTTCGGT AGCCAGAGCT AGCCAGAGCT	IFLRQEIDRM FSTLGFWFTE MVLCNEVTKW AENNTLRDPR ID NO:47 PDG5 to start and stop of 41 CAGGGGTTGC TCCAGARAAG GATCCAGACA TCTCTCTCAC	QRVLSLVRST LIERNSQFTS MKDDISTPPT FYSCPIYKKP DNA SEQUENCE odons) 51 TTTCAGATAA CATCAGCAGC CATACCATGA TGATGGTGGA	180 240 300 360 60 120 180 240
45 50	TREAVVARLA LTELKLAIDG WVFNGRPHCF EGYVYVGLYL VRTDLNYIAA Nucleic Acid Acce Coding sequence: 1 GGAGCAGCCT AGATGACATG ACAGCCCATA AGATGCAGCT AAGCCTTTCT	DDMLEKLPPD TIIMSENLQD WMTGFFNPQG EGAGWDERMM VDLRTAQTPE SSION #: AB033 68-3345 11 ACAACTTCAC GGAAGGAGAA CCTGAAAACA TCTGGAGCTG ACACCCAAG	YVPFEVKERL ALDCMFDARI FLTAMRQEIT KLIESKPKVL HWVLRGVA 036 (underlined sequ 21 AACCAGAAAC ATGCTGGCAI TGGACAATTC AGAAGACCAG AGAGGCCCAI	OKMGPFOPMN PAWWKKASWV RANKGWALDN FELMPVIRTY LL CDVK SEC ences correspond 31	IFLRQEIDRM FSTLGFWFTE MVLCNEVTKW AENNTLRDPR 2 ID NO:47 PDG5 to start and stop of 41 CAGGGGTTGC TCCAGAAAAG GATCCACACA TCTCTCTCACC GCAGCAGGG GCAGCAGGG	QRVLSLVRST LIERNSQFTS MKDDISTPPT FYSCPIYKKP DNA SEQUENCE odons) 51 TTTCAGATAA CATCAGCAGC CATACCATGA TGATGGTGGA TCATGGTGGT	180 240 300 360 60 120 . 180 240 300
45 50	TREAVVARLA LTELKLAIDG WVFNGRPHCF EGYVYVGLYL VRTDLNYIAA Nucleic Acid Acce Coding sequence: 1 GGAGCAGCCT AGATGACATG ACAGCCCATA AGATGCAGCT AAGCCTTTCT	DDMLEKLPPD TIIMSENLQD WMTGFFNPQG EGAGWDERMM VDLRTAQTPE SSION #: AB033 68-3345 11 ACAACTTCAC GGAAGGAGAA CCTGAAAACA TCTGGAGCTG ACACCCAAG	YVPFEVKERL ALDCMFDARI FLTAMRQEIT KLIESKPKVL HWVLRGVA 036 (underlined sequ 21 AACCAGAAAC ATGCTGGCAI TGGACAATTC AGAAGACCAG AGAGGCCCAI	OKMGPFOPMN PAWWKKASWV RANKGWALDN FELMPVIRTY LL CDVK SEC ences correspond 31	IFLRQEIDRM FSTLGFWFTE MVLCNEVTKW AENNTLRDPR 2 ID NO:47 PDG5 to start and stop of 41 CAGGGGTTGC TCCAGAAAAG GATCCACACA TCTCTCTCACC GCAGCAGGG GCAGCAGGG	QRVLSLVRST LIERNSQFTS MKDDISTPPT FYSCPIYKKP DNA SEQUENCE odons) 51 TTTCAGATAA CATCAGCAGC CATACCATGA TGATGGTGGA	180 240 300 360 60 120 180 240
45 50 55	TREAVVARLA LTELKLAIDG WVFNGRPHCF EGYVYVGLYL VRTDLNYIAA Nucleic Acid Acce Coding sequence: 1 GGAGCAGCCT AGATGACATG ACAGCCCATA AGATGCAGCT TATGAATCCT	DDMLEKLPPD TIIMSENLQD WMTGFFNPQG EGAGWDERNM VDLRTAQTPE SSIon #: AB033 68-3345 11 ACAACTTCAC GGAAGGAGAA CCTGAAAACA TCTGGAGCTG ACAACCCAAG	YVPFEVKERL ALDCMFDARI FLTAMRQEIT KLIESKPKVL E HWVLRGVA 036 0 (underlined sequ 21 AACCAGAAAAC ATGCTGGCAT TGGACAATTC AGAAGACAGA AGGAGGCCAT AGGAGGCCAT AGTTAGAAGA	OKMGPFOPMN PAWWKKASWV RANKGWALDN FELMPVIRIY LL CDVK SEC ences correspond 31	IFLRQEIDRM FSTLGFWFTE MVLCNEVTKW AENNTLRDPR 2 ID NO:47 PDG5 to start and stop of 41 CAGGGGTTGC TCCAGAAAAG GATCCACACA TCTCTCTCACC GCAGCAGGG GCAGCAGGG	QRVLSLVRST LIERNSQFTS MKDDISTPPT FYSCPIYKKP DNA SEQUENCE odons) 51 TTTCAGATAA CATCAGCAGC CATACCATGA TGATGGTGGA TCTTGGTGGA ATTTACAAAA	180 240 300 360 60 120 . 180 240 300
45 50 55	TREAVVARLA LTELKLAIDG WVFNGRPHCF EGYVYYGLYL VRTDLNYIAA Nucleic Acid Acce Coding sequence: 1 GAGGCAGCT ACAGCCCATA AGATGCAGT AAGCCTTTCT TATGAATCC GGCCCAATACC	DDMLEKLPPD TIIMSENLQD WMTGFFNPQG EGAGWDKRNM VDLRTAQTFF SSION #: AB033 68-3349 11 ACAACTTCAC GGAAGGAGAA CCTGAAAACA TCTGGAGCTG ACAACCCAAG TCTCATATCC AAAATGGAGT	YVPFEVKERL ALDCMFDARI FLTAMRQEIT KLIESKPKVL E HWVLRGVA 036 0 (underlined sequence) AACCAGAAAA ATGCTGGCAT TGGACAATTC AGAAGACAGA AGGAGGCCAT AGTAGAAGA AGTAGAAGA AGTAGAAGA CAGCCCAGGA	QKMGPFQPMN PAWWKKASWV RANKGWALDN FELMPVIRTY LL CDVK SEC ences correspond 31 CACTACCCCT AGATTTCGGA CATGGTTAGT AGCCAGAGCT TCTCAGTA TCAAGAAGCT TCATCAACTA TCAAGAAGCT	IFLRQEIDRM FSTLGFWFTE WVLCNEVTKW AENNTLRDPR ID NO:47 PDG5 to start and stop of 41 CAGGGGTTGC TCCAGAAAAG GATCCACAAC TCTCTCTCAC GCAGCAGAGG TTCAGCTTTG ATCTGCAAAG ATCTGCAAAAG	QRVLSLVRST LIERNSQFTS MKKDDISTPPT FYSCPIYKKP DNA SEQUENCE odons) 51 TITCAGATAA CATCAGCAGC CATACCATGA TGATGGTGGA TGATGGTGGA CTCAGGTGTT ATTTACAAAA AAAAGCCTTC	60 120 180 240 300 360
45 50	TREAVVARLA LTELKLAIDG WVFNGRPHCF EGYVYYGLYL VRTDLNYIAA Nucleic Acid Acce Coding sequence: 1 GGAGCAGCCT AGATGACATG ACAGCCATA AGATGCAGCT TATGAATCC TGGCCAATCC TGGAAATGTT	DDMLEKLPPD TIIMSENLQD WMTGPFNPQG EGAGWDKRMM VDLRTAQTPF SSION #: AB033 68-3345 11 ACAACTTCAC GGAAGGAGAA TCTGGAGCTG ACAACCCAAG TCTCATATCC ACAATGGAGTG ACAATGGAGTG ACAATGGAGTG ACAATGGAGTG ACAATGGAGTG ACACCAGACCT CACCAGACCT	YVPFEVKERL ALDCMFDARI FLTAMRQEIT FLTAMRQEIT KLIESKPKVL E HWVLRGVA 036 Qunderlined sequ 21 AACCAGAAAC ATGCTGGCAT TGGACAATTC AGAAGACAGA AGGAGGCCAI AGTTAGAAG CAGCCCAGGA CTTACAGCAAG	QKMGPFQPMN PAWWKKASWV RANKGWALDN FELMPVIRTY LL CDVK SEC ences correspond 31 CACTACCCTI AGATTTCGGT AGCCAGAGCT TCTCAGTAGT TGTTCAAGAACT TGTTTTGGGT	IFLRQEIDRM FSTLGFWFTE MVLCNEVTKW AENNTLRDPR ID NO:47 PDG5 to start and stop of 41 CAGGGGTTGC TCCAGARAGG GATCACAAC TCTCTCTCAC GCAGCAGAGG TTCAGCTTGA ATCTGCTAAA ATGACAAGTA	QRVLSLVRST LIERNSQFTS MKXDDISTPPT FYSCPIYKKP DNA SEQUENCE DOORS) 51 TTTCAGATAA CATCAGCAGC CATACCATGA TGATGGTGGA CTCAGGTGTT ATTTACARAA AAAAGCCTTC CTACAGCCAA	180 240 300 360 60 120 240 300 360 420 480
45 50 55	TREAVVARLA LTELKLAIDG WVFNGRPHCF EGYVYVGLYL VRTDLNYIAA Nucleic Acid Acce Coding sequence: 1 GGAGCAGCCT AGATGACATG ACAGCCCATA AGATGACATC TATGAATCCT GGCCAATCCT TGGAAATGTT AGAGAGTTTTT AGGAGATGTTT	DDMLEKLPPD TIIMSENLQD WMTGFFNPQG EGAGWDERMM VDLRTAQTPE SSION #: AB033 68-3345 11 ACAACTTCAC GGAAGGAGAA CCTGAAACA TCTGGAGCTG ACAACCCAAG TCTCATATCC AAAATGGAGT CACCAGACCT TATGCCAAGA	YVPFEVKERL ALDCMFDARI FLTAMRQEIT KLIESKPKVL HWVLRGVA 036 Qunderlined sequ 21 AACCAGAAAA ATGCTGGCAT AGGAGACCA AGGAGGCCAT AGTTAGAAGA CAGCCCAGGA TTACAGCAAG CTCTGCCTCC	OKMGPFOPMN PAWWKKASWV RANKGWALDN FELMPVIRTY LL CDVK SEC ences correspond 31	IFLRQEIDRM FSTLGFWFTE MVLCNEVTKW AENNTLRDPR ID NO:47 PDG5 to start and stop of 41 CAGGGGTTGC TCCAGAAAAG GATCACAAG GCAGCAGAGG TTCAGCTTG ATCTGCAAAG ATGACAAGTA TTTCAGTCCT	QRVLSLVRST LIERNSQFTS MKDDISTPPT FYSCPIYKKP DNA SEQUENCE odons) 51 TTTCAGATAA CATCAGCAGC CATACCATGA TGATGGTGGA TGATGGTGGT ATTTACAAAA AAAAGCCTTC CTACAGCAGC CAAGGAAGCC CAAGGAAGCC	60 120 120 120 1300 360 420 480 540
45 50 55	TREAVVARLA LTELKLAIDG WVFNGRPHCF EGYVYVGLYL VRTDLNYIAA Nucleic Acid Acce Coding sequence: 1 GGAGCAGCCT AGATGACATG ACAGCCCATA AGATGCAGCT TATGAATCCT GGCCCAATCC TGGAAATGT TGGAGAGTT TGATGCTGAA	DDMLEKLPPD TIIMSENLQD WMTGFFNPQG EGAGWDKRNM VDLRTAQTPE SSION #: AB033 68-3345 11 ACAACTTCAC GGAAGGAGAA CCTGAAAACA TCTGGAGCT ACAACCCAAGC TCTCATATCC AAAATGGAGT TATGCCAAGA GAAGTCTCCT	YVPFEVKERL ALDCMFDARI FLTAMRQEIT KLIESKPKVL E HWVLRGVA 036 0 (underlined sequ 21 AACCAGAAAAC ATGCTGGCAT TGGACAATTC AGAAGACAGA AGGAGGCCAT AGTTAGAAGA CTCTGCCTCC CAGATTCAGA	OKMGPFOPMN PAWWKKASWV RANKGWALDN FELMPVIRIY LL CDVK SEC ences correspond 31	IFLRQEIDRM FSTLGFWFTE MVLCNEVTKW AENNTLRDPR 2 ID NO:47 PDG5 to start and stop of 41 CAGGGGTTGC TCCAGAAAAG GATCCACAAC GCAGCAGAGG TTCAGCTTTG ATCTGCCAAAA ATTTCAGTCCT GAGGAGGGGGGGGGG	QRVLSLVRST LIERNSQFTS MKDDISTPPT FYSCPIYKKP DNA SEQUENCE odons) 51 TTTCAGATAA CATCAGCAGC CATACCATGA TGATGGTGGA TGATGGTGGA CTCAGGCGTT ATTTACAAAA AAAAGCCTTC CTACAGCCAGC ATGGTAGCAAGCC ATGGTAGCAAGCCAAGGAAGACC ATGGTACCATGA	60 120 180 240 300 360 120 180 240 420 420 480 540 600
45 50 55	TREAVVARLA LTELKLAIDG WVFNGRPHCF EGYVYYGLYL VRTDLNYIAA Nucleic Acid Acce Coding sequence: GGAGCAGCCCTA AGATGACATG AGATGCAGCT TATGAATCCT TGGAAATGTT AGGAGATGTT AGAGCTGCT TGAAACTGGCT TGAACTGGCT AGGAGATGTT AGAGCTGCT AGGAGATGTT AGAACTGGCT	DDMLEKLPPD TIIMSENLQD WMTGPFNPQG EGAGWDKRNM VDLRTAQTPF SSION #: AB033 68-3345 11 ACAACTTCAC GGAAGGAGAA CCTGAAAACA TCTGGAGCTG ACAACCCAAGA TCTCATATCC AAAATGGAGT CACCAGACCT TATGCCAAGA GAAGTCTCCT CATGGTCACT	YVPFEVKERL ALDCMFDARI FLTAMRQEIT KLIESKPKVL E HWVLRGVA 036 0 (underlined sequ 21 AACCAGAAAA ATGCTGGCAT TGGACAATTC AGAAGACAGA AGGAGGCCAT AGTTAGAAGA CAGCCAGGA TTACAGCAAG CTTCCCAGTC CAGATTCAGA CTTCCCAGTC	QKMGPFQPMN PAWWKKASWV RANKGWALDN FELMPVIRTY LL CDVK SEC ences correspond 31	IFLRQEIDRM FSTLGFWFTE WVLCNEVTKW AENNTLRDPR ID NO:47 PDG5 to start and stop of 41 CAGGGGTTGC TCCAGAAAAG GATCACAAAC TCTCTCTCAC GCAGCAGAGG TTCAGCTTTG ATCTCCAAAG ATGACAAGTA TTTCAGTCCT TGAGGAGGGG TTTGAGGAGGG TTTGAGGAGGG	QRVLSLVRST LIERNSQFTS MKKDDISTPPT FYSCPIYKKP DNA SEQUENCE Ddons) 51 TTTCAGATAA CATCAGCAGC CATACCATGA TGATGGTGGA CTCAGGTGTA ATTTACAAAT AATAGCCTTC CTACAGCCAA CAAGGAAGCC CATGGTTCTGA AACAAGAAGT	180 240 300 360 60 120 180 240 300 360 420 480 540 660
45 50 55 60	TREAVVARLA LTELKLAIDG WVFNGRPHCF EGYVYYGLYL VRTDLNYIAA Nucleic Acid Acce Coding sequence: 1 GGAGCAGCCT AGATGACATG ACAGCCCATG AGACCTTCT TATGAATCCT GGCCCAATCC GGCCAATCT TGGAAATGTT TGGAGATGTT TGGATGCTGAGA	DDMLEKLPPD TIIMSENLQD WMTGFFNPQG EGAGWDKRMM VDLRTAQTFF SSION #: AB033 68-3345 11 ACAACTTCAC GGAAGGAGAA TCTGGAGCTG ACAACCCAAG TCTCATTATCC AAAATGGAGCT TATGCCAAGA GAAGTCTCCT TATGCCAAGA CATGGTCACT TCATAGGTCACT TCATGGTCACT TTATGCTAAGA	YVPFEVKERL ALDCMFDARI FLTAMRQEIT KLIESKPKVL E HWVLRGVA 036 Qunderlined sequence AACCAGAAAAA ATGCTGGCAT AGGAGACAATA AGGAGGCCAGA AGGTTAGAAGA CAGCCAGGAA TTACAGCAGA CTCTGCCTCC CAGATTCAGAGA CTTCCCAGTC TTGTTGAGGGA	QKMGPFQPMN PAWWKKASWV RANKGWALDN FELMPVIRTY LL CDVK SEC ences correspond 31	IFLRQEIDRM FSTLGFWFTE WVLCNEVTKW AENNTLRDPR ID NO:47 PDG5 to start and stop of CAGGGGGTTGC TCCAGAAAAG GATCACAAA TCTCTCTCAG GCAGCAGAGG TTCAGCTTAG ATGACAAGTA TTTCAGTCCT GAGGAGGGGGG GTTGAAGATC TCTGAAGAAGT TTTGAAGATG TCTGAAGAGT TCTGAAGAGT TCTGAAGAGGGGGGGGGG	QRVLSLVRST LIERNSQFTS MKDDISTPPT FYSCPIYKKP DNA SEQUENCE DOORS) 51 TTTCAGATAA CATCAGCAGC CATACCATGA TGATGGTGGA CTCAGGTGTT ATTTACABAA AAAGCCTTC CTACAGCCAA CAAGGAAGCC ATGGTTCTGA AACAAGAAGT AGCTGGACCT	180 240 300 360 60 120 - 180 240 300 360 420 540 600 660 720
45 50 55	TREAVVARLA LTELKLAIDG WVFNGRPHCF EGYVYYGLYL VRTDLNYIAA Nucleic Acid Acce Coding sequence: 1 GGAGCAGCCT AGATGACATG ACAGCCCATG AGACCTTCT TATGAATCCT GGCCCAATCC GGCCAATCT TGGAAATGTT TGGAGATGTT TGGATGCTGAGA	DDMLEKLPPD TIIMSENLQD WMTGFFNPQG EGAGWDKRMM VDLRTAQTFF SSION #: AB033 68-3345 11 ACAACTTCAC GGAAGGAGAA TCTGGAGCTG ACAACCCAAG TCTCATTATCC AAAATGGAGCT TATGCCAAGA GAAGTCTCCT TATGCCAAGA CATGGTCACT TCATAGGTCACT TCATGGTCACT TTATGCTAAGA	YVPFEVKERL ALDCMFDARI FLTAMRQEIT KLIESKPKVL E HWVLRGVA 036 Qunderlined sequence AACCAGAAAAA ATGCTGGCAT AGGAGACAATA AGGAGGCCAGA AGGTTAGAAGA CAGCCAGGAA TTACAGCAGA CTCTGCCTCC CAGATTCAGAGA CTTCCCAGTC TTGTTGAGGGA	QKMGPFQPMN PAWWKKASWV RANKGWALDN FELMPVIRTY LL CDVK SEC ences correspond 31	IFLRQEIDRM FSTLGFWFTE WVLCNEVTKW AENNTLRDPR ID NO:47 PDG5 to start and stop of CAGGGGGTTGC TCCAGAAAAG GATCACAAA TCTCTCTCAG GCAGCAGAGG TTCAGCTTAG ATGACAAGTA TTTCAGTCCT GAGGAGGGGGG GTTGAAGATC TCTGAAGAAGT TTTGAAGATG TCTGAAGAGT TCTGAAGAGT TCTGAAGAGGGGGGGGGG	QRVLSLVRST LIERNSQFTS MKKDDISTPPT FYSCPIYKKP DNA SEQUENCE Ddons) 51 TTTCAGATAA CATCAGCAGC CATACCATGA TGATGGTGGA CTCAGGTGTA ATTTACAAAT AATAGCCTTC CTACAGCCAA CAAGGAAGCC CATGGTTCTGA AACAAGAAGT	180 240 300 360 60 120 - 180 240 300 360 420 540 600 660 720
45 50 55 60	TREAVVARLA LTELKLAIDG WVFNGRPHCF EGYVYVGLYL VRTDLNYIAA Nucleic Acid Acce Coding sequence: 1 GGAGCAGCCT AGATGACATG ACAGCCCATA AGATGACATG TATGAATCCT TGGAAATGTT TGGAGATGTT TGATGCTGAA AGACTGCT CTTCTCAGAA CAGATGCCTC	DDMLEKLPPD TIIMSENLQD WMTGFFNPQG EGAGWDERMM VDLRTAQTPE SSION #: AB033 68-3345 11 ACAACTTCAC GGAAGGAGAA CCTGAAAAC TCTGGAGCTG ACAACTCAGA CTTATATCC AAAATGGAGCT TCACAGACCT TATGCCAAGA GAAGTCTCCT TCACAGACCT TCAC	YVPFEVKERL ALDCMFDARI FLTAMRQEIT KLIESKPKVL HWVLRGVA 036 Qunderlined sequ 21 AACCAGAAAA ATGCTGGCAT AGAAGACAAT AGGAGGCCAGA AGGAGGCCAGA AGTTAGAAGA CAGCCCAGGA TTACAGCAAG CTCTGCCTCC CAGATTCAGAG TTGCTAGAGGA TTGCTAGAGGA TTGCTAGAGGA TTGCTAGAGGAGC TTGTTGAGGGA	QKMGPFQPMN PAWWKKASWV RANKGWALDN FELMPVIRTY LL CDVK SEC ences correspond 31	IFLRQEIDRM FSTLGFWFTE MVLCNEVTKW AENNTLRDPR ID NO:47 PDG5 to start and stop of 41	QRVLSLVRST LIERNSQFTS MKDDISTPPT FYSCPIYKKP DNA SEQUENCE DOORS) 51 TTTCAGATAA CATCAGCAGC CATACCATGA TGATGGTGGA CTCAGGTGTT ATTTACABAA AAAGCCTTC CTACAGCCAA CAAGGAAGCC ATGGTTCTGA AACAAGAAGT AGCTGGACCT	180 240 300 360 60 120 - 180 240 300 360 420 600 600 600 720 780
45 50 55 60	TREAVVARLA LTELKLAIDG WVFNGRPHCF EGYVYYGLYL VRTDLNYIAA Nucleic Acid Acce Coding sequence: 1 GGAGCAGCCT AGATGACATG ACAGCCCATA AGATGCAGT TATGAATCCT GGCCCAATCC TGGAATGTT TGATGCTGAA AGACTGCT TTGATGCTGAA AGACTGCCT CTTCTCAGAA CAGATGCCTC CAGTTATGTT	DDMLEKLPPD TIIMSENLQD WMTGFFNPQG EGAGWDKRNM VDLRTAQTPF SSION #: AB033 68-3349 11 ACAACTTCAC GGAAGGAGAA CCTGAAAACA TCTGGAGCTG ACAACCCAGACTT TATGCCAAGA GAAGTCTCCT TATGCCAAGA CATGGTCACT TCCAAGACTT TCCCAGGCTT TCCCAGGCTT GAAAAGTTACCAGACTC GAAAAGTTACCAGACTATCCAAAAGTT TCCCAGGCTT GAAAAGTTACCAGCTT GAAAAGTTACA	YVPFEVKERL ALDCMFDARI FLTAMRQEIT KLIESKPKVL E HWVLRGVA) 036 0 (underlined sequence of the common co	QKMGPFQPMN PAWWKKASWV RANKGWALDN FELMPVIRTY LL CDVK SEC ences correspond 31 CACTACCCCT AGATTTCGGA CATGGTTAGT AGCAGAGCT TCTCTCAGT AGAAGATC TGTTTAGGA CTTGAGAGCT CAGAAGCT CAGAAGATGCA CTTGAGCAGC TGAAGATGCA TGATTGCAGC	IFLRQEIDRM FSTLGFWFTE WVLCNEVTKW AENNTLRDPR ID NO:47 PDG5 to start and stop of 41 CAGGGGTTGC TCCAGAAAG GATCCACAAC TCTCTCACA GCAGCAGAGG ATGACAAGA ATGACAAGA ATGACAAGT TTCAGTCCT GAGGAGGGGG TTTGAGGAGG TTTGAGGAGG GAAGTCTTCA AGCTCTAGAGGAG	QRVLSLVRST LIERNSQFTS MKDDISTPPT FYSCPIYKKP DNA SEQUENCE odons) 51 TTTCAGATAA CATCAGCAGC CATACCATGA TGATGGTGGA CTCAGGTGTT ATTTACAAAA AAAGCCTTC CTACAGCAAA CAAGGAAGCC ATGGTTCTGA AACAAGAAGC AGGTGAACCT CAGGAATCAAG CAGGAACCAT CAGGAATCAAG	180 240 300 360 60 120 180 240 360 420 480 540 660 720 780 840
45 50 55 60	TREAVVARLA LTELKLAIDG WVFNGRPHCF EGYVYYGLYL VRTDLNYIAA Nucleic Acid Acce Coding sequence: 1 GGAGCAGCATT AGATGACATT AGATGACATT TATGAATCCT TGGAGATGTT AGAGGTTT TGGAGATGTT AGAGGTTT TGGAGATGTT AGAGTGTT AGAGTGTT AGAGTGTT AGAGTGTT AGAGTGTT TGAGATGCTC CTTCTCAGAA CAGTTATGTT TCTCAGACAC	DDMLEKLPPD TIIMSENLQD WMTGPFNPQG EGAGWDKRNM VDLRTAQTPF SSION #: AB033 68-3345 11 ACAACTTCAC GGAAGGAGA CCTGAAAACA TCTGGAGCTG ACAACCCAAG TCTCATACC AAAATGGAGT CACCAGACCT TATGCCAAGA GAAGTCTCCT CATGGTCACT TCCAGGCTT TCCAGGCTT TCCAGGCTT TCCAGGCTT TCCAGGCTCACC CAAAAGTACA CCTGCTCAGGC CCTGCTCAGGC CCTGCTCAGGC CCTGCTCAGGC CCTGCTCAGGC CCTGCTCAGGC TGAAAAGTACA CCTGCTCAGGC CCTGCTCAGC CCTGCT	YVPFEVKERL ALDCMFDARI FLTAMRQEIT KLIESKPKVL E HWVLRGVA 036 Qunderlined sequ 21 AACCAGAAAAC ATGCTGGCAA TGGACAATTC AGAAGACAGA AGGAGGCCAA AGGAGGCCAA AGTTAGAAG CTTCTGCTCC CCAGTTCAGA CTTCTCAGAC TTGTTGAGGAC TAGAGGAGGCC ACACTTCTGA	QKMGPFQPMN PAWWKKASWV RANKGWALDN FELMPVIRTY LL CDVK SEC ences correspond 31	IFLRQEIDRM FSTLGFWFTE WVLCNEVTKW AENNTLRDPR ID NO:47 PDG5 to start and stop of 1 CAGGGGTTGC TCCAGAAAG GATCCACAAG TCTCTCTCAC GCAGCAGAGG TTCAGCTTG TTCAGCTTG TTCAGCTTG TTCAGCTG TTCAGAGAGG ATGAGAGGG TTTGAGGAGGG TTTGAGGAGG TTTGAGGAGG TTTGAGGAGG GAAGTCTTCA AGCTCAGAGG CAACAAGAAG	QRVLSLVRST LIERNSQFTS MKXDDISTPPT FYSCPIYKKP DNA SEQUENCE odons) 51 TTTCAGATAA CATCAGCAGC CATACCATGA TGATGGTGGA CTCAGGTGTT ATTTACAAGA AAAAGCCTTC CTACAGCCAA CAAGGAAGCC ATGGTTCTGA AACAAGAAGT AGCTGGACCT CAGAATCAAG AAGACTGCC TCTCCTCTGC	180 240 300 360 60 120 180 240 300 360 540 660 720 780 840 900
45 50 55 60	TREAVVARLA LTELKLAIDG WVFNGRPHCF EGYVYYGLYL VRTDLNYIAA Nucleic Acid Acce Coding sequence: 1 GGAGCAGCCT AGATGACATG ACAGCCCATG AGACTCT TATGAATCATC GGCCCAATCT TGGAAATCTT TGGAGATGTT TGATGCTGAA AGACTGCT CTTCTCAGAA CAGATGCCTC CAGTTATTT TCTCAGAACAC TTCCAGACAC TTCCAGACAC TTCCAGACAC TTCCAGACAC	DDMLEKLPPD TIIMSENLQD WMTGFFNPQG EGAGWDKRMM VDLRTAQTPF SSION #: AB033 68-3345 11 ACAACTTCAC GGAAGGAGAA TCTGGAGCTG ACAACCCAAG TCTGATCAC AAATGGACT TATGCCAAGA GAAGTTCACT TCAGAGCTT TCACAGACCT TTAGAAAAAT TCCCAGGCTT TCAGAAAAAAA AATGGACT TCATGCTCAGG AAAATTACAC AAAAAAAAAA	YVPFEVKERL ALDCMFDARI FLTAMRQEIT FLTAMRQEIT KLIESKPKVL E HWVLRGVA 036 Qunderlined sequence AACCAGAAAAA ATGCTGGCAT AGGAGACAAT AGGAGGCCAGAA AGGAGGCCAGAA CAGCCCAGGA TTACAGCAAA CTTCCCAGT TTGTTGAGGAA TTGTTGAGGAA TTGTTGAGGAA CTTCCCAGT TTGTTGAGGAA CTTCCCAGT TTGTTGAGGAA CTTCCCAGT TTGTTGAGGAA CTTCCCAGT CTTGCTGAGAAAAAAAAAA	QKMGPFQPMN PAWWKKASWV RANKGWALDN FELMPVIRTY LL CDVK SEC ences correspond 31	IFLRQEIDRM FSTLGFWFTE WVLCNEVTKW AENNTLRDPR ID NO:47 PDG5 to start and stop of 41 CAGGGGTTGC TCAGARARG GCATCACARG TCTCTCTCAC GCAGCAGAGG TTCAGCTTG ATTCAGCTTG GAGGAGGGGG GTTGAGGAGG TTTGAAGATA TTTCAGTCCT CAGGAGGGGGG GAGGAGGAGGGGGG GAGGAGAGAG GAGGTCTTCA AGCTCAGAGAG CAGCTAGCAGAG CAGCTGCCTT	QRVLSLVRST LIERNSQFTS MKDDISTPPT FYSCPIYKKP DNA SEQUENCE DOORS) 51 TTTCAGATAA CATCAGCAGC CATACCATGA TGATGGTGGA CTCAGGTGTT ATTTACABAA AAAAGCCTTC CTACAGCCAA CAAGGAAGCC ATGGTTCTGA AACAAGAAGT AACAAGAAGT AACAAGAAGT CAGAATCAAG AAGACCTGCC TCTCCTCTGC CCCAGATGCCC	180 240 300 360 60 120 - 180 240 300 360 420 660 720 780 840 900 960
45 50 55 60	TREAVVARLA LTELKLAIDG WVFNGRPHCF EGYVYVGLYL VRTDLNYIAA Nucleic Acid Acce Coding sequence: 1 GGAGCAGCCTT AGATGACATG ACAGCCCATA AGACTCTTCT TATGAATCCT TGGAAATGTT TGATGCTGAA AGACTGCTCC CTTCTACAGAA CAGATGCTT CTTCTCAGAA TTCTCAGACAC TTCAAATAAT TTCTCAGCCC	DDMLEKLPPD TIIMSENLQD WMTGFFNPQG EGAGWDERMM VDLRTAQTPE SSION #: AB033 68-3345 11 ACAACTTCAC GGAAGGAGAA CCTGAAAAC TCTGGAGCTG ACAACTCAGG TCTCATATCC AAAATGGAGC TATGCCAAGA GAAGTCTCCT TCAAAAAGTT TCCCAGGCTT TCAAAAAGTT TCCCAGGCTT TCAAAAAGTT TCCCAGGCTT TCAAAAAGTT TCCCAGGCTT TCAAAAAGTT TCCCAGGCTT TCAAAAAGTT TCCCAGGCTT ACACGGCTT TCAAAAAGTT TCCCAGGCTT ACACGGCTT TCAAAAAGTT CCTGCTCAGG ACTCCTGAAG	YVPFEVKERL ALDCMFDARI FLTAMRQEIT KLIESKPKVL HWVLRGVA 036 Qunderlined sequ 21 AACCAGAAAA ATGCTGGCAT AGAAGACAAT AGATAGAAGACAAT AGATAGAAGACAAA CTCTGCCTCC CAGATTCAGAA TTGCAGAA TTGCAGAA CTTCTCAGAC TTGTTGAGGA TAGAGGAGCC ACATTCTGA AGACGAAAG CTCTCCCAGT CTTGTTGAGGA TAGAGGAGCC ACATTCTGA AGCAGAATGA AGCAGAATGA CTTGTTGAGGA AGCAGAATGA CTACTGTTCA	QKMGPFQPMN PAWWKKASWV RANKGWALDN FELMPVIRTY LL CDVK SEC ences correspond 31	IFLRQEIDRM FSTLGFWFTE MVLCNEVTKW AENNTLRDPR ID NO:47 PDG5 to start and stop of 41	QRVLSLVRST LIERNSQFTS MKDDISTPPT FYSCPIYKKP DNA SEQUENCE odons) 51 TTTCAGATAA CATCAGCAGC CATACCATGA TGATGGTGGA CTCAGGTGTT ATTTACAAAA AAAGCCTTC CTACAGCACA CAAGGAAGCC ATGGTTCTGA AACAAGAAGT CAGAATCAAG AAGACCTGCC CCAGATTGCC CCAGTTGCC CCAGTTGCC CCAGTTGCCC CAGTTGCC CCAGTTGCC CCAGTTGCC CCAGTTGCC CCAGTTGCC CCAGTTGGCCA	180 240 300 360 60 120 120 240 300 360 420 600 660 720 780 840 900 960 1020
45 50 55 60	TREAVVARLA LTELKLAIDG WVFNGRPHCF EGYVYYGLYL VRTDLNYIAA Nucleic Acid Acce Coding sequence: 1 GGAGCAGCCT AGATGACATG AGATGCATG AGATGCATCT TATGAATCCT TGGAAATGT AGACTGTAA AGACTGCTT CATGATGCTT CATGATCT TCATGATCT TCTCAGACA CAGATGCTT TCTCAGACAC TTCTAAGATACT TCTCAGACAC TTCTAAGATACT TTCTCAGCAC TTCTAATAAA	DDMLEKLPPD TIIMSENLQD WMTGFFNPQG EGAGWDKRMM VDLRTAQTPE SSION #: AB033 68-3349 11 ACAACTTCAC GGAAGGAGAA CCTGAAAACA TCTGGAGCTG ACAACCCAAGA TCTCATATCC AAAATGGAGTT TCCCAGGACTT TCCCAGGACTT TCCCAGGACTT TCCCAGGACTT TCCCAGGACTT TCCCAGGACTT TCCCAGGACTT TCCCAGGACTT TCCCAGGACTT TCCAGGACTT TCCAGGACTT TCCAGGACTT TCAAAAAGTTACA CCTGCTCAAGA ATTATGAATC CAGAGCGATT	YVPFEVKERL ALDCMFDARI FLTAMRQEIT FLTAMRQEIT KLIESKPKVL E HWVLRGVA 036 0 (underlined sequ 21 AACCAGAAAC ATGCTGGCAT TGGACAATTC AGAAGACAGA AGAGCAGAT AGTTAGAGA CTTCTGCCTC CAGATTCAGA CTTCTTAGGAGC AGAGTTCAGA CTTCTTGTTGAGGA CTTCTTGTTGAGGA CTTCTTGTTGAGGA CTTCTTGAGGA CTTCTTGAGGA CTTCTTGAGGA CTTCTGGGAAA CCTTCTGGCAAA CCTTCTGGCAAA CCTTCTGGCAAA CCTTCTGGAAAA CCTTCTGGAAAA CCTTCTGGAAAA CCTTCTGGAAAA CCTTCTGGAAAA CCTTCTGGAAAC CCTTGGGAAAC CCTAGTGGAGCC	QKMGPFQPMN PAWWKKASWV RANKGWALDN FELMPVIRTY LL CDVK SEC ences correspond 31	IFLRQEIDRM FSTLGFWFTE WVLCNEVTKW AENNTLRDPR ID NO:47 PDG5 to start and stop of 41 CAGGGGTTGC TCCAGAAAG GATCCACAAC TCTCTCTCAC GCAGCAGAGG TTCAGCTTTG ATGACAGTA TTTCAGTCTT GAGGAGGGGG TTTGAAGATG TCTGAGGAGG GAACTCTTCA AGCTCAGAGG CAACAAGAAG CAGCTGCCTT AGACACCCTT AGACACCCTT	QRVLSLVRST LIERNSOFTS MKDDISTPPT FYSCPIYKKP DNA SEQUENCE dons) 51 TTTCAGATAA CATCAGCAGC CATACCATGA TGATGGTGGA CTCAGGTGTT CTACAGCAAA AAAAGCCTTC CTACAGCAAA AACAAGAAGT AGCTGGACCT CAGAATCAAG AACAACCAGC TCTCTCTCC CCAGTGGCCC TCTCCTCTCC	60 120 120 120 120 140 300 360 420 480 540 660 720 780 960 1020 1080
45 50 55 60	TREAVVARLA LTELKLAIDG WVFNGRPHCF EGYVYYGLYL VRTDLNYIAA Nucleic Acid Acce Coding sequence: 1 GGAGCAGCCATA AGATGCACTT AAGACCCATA AGACCCATA AGAGCTTTCT TATGAATCCT TGGACATGTT AGAGGTTTAT AGAGTGCTC CAGATGCTC CAGATGCTC CAGTTATTT TCTCAGCAC TTCTAGAACAC TTCTAGAACAC TTCTAGAACAC TTCTAGAACAC TTCTAGAACAC TTCTAGAACAC GGTGAACCCT TCTATAAAAA GGTGAACCCT	DDMLEKLPPD TIIMSENLQD WMTGPFNPQG EGAGWDKRMM VDLRTAQTPE ssion #: AB033 68-3349 11 ACAACTTCAC GGAAGGAGAA CCTGAAAACA TCTGGAGCTG ACAACCCAAG TCTCATATCCAC CACAGACCT TATGCCAAGA GAAGTCTCCT TCAAAAAGTT TCCCAGGCTT TCCAGGCTT TCCAAGACT CATGGTCACT CAAAAGTT CCCAGGCTT TCCAAGACT CATGGTCACT CAAAAGTT CCCAGGCTT CAAAAGTT CCCAGGCTT CAAAAGTT CCCAGGCTT AAAGTCACG ACTCCTGAAG ATTATGAAT AAAGTGGAGC	YVPFEVKERL ALDCMFDARI FLTAMRQEIT FLTAMRQEIT FLTEMRYVL E HWVLRGVA 036 Qunderlined sequ 21	QKMGPFQPMN PAWWKKASWV RANKGWALDN FELMPVIRTY LL CDVK SEC ences correspond 31	IFLRQEIDRM FSTLGFWFTE WVLCNEVTKW AENNTLRDPR ID NO:47 PDG5 to start and stop of the start and star	QRVLSLVRST LIERNSQFTS MKXDDISTPPT FYSCPIYKKP DNA SEQUENCE odons) 51 TTTCAGATAA CATCAGCAGC CATACCATGA TGATGGTGGA CTCAGGTGTT ATTTACAAGA AAAAGCCTTC CTACAGCAA AACAAGCAGC ATGGTTCTGA AACAAGAAGT AGCTGGACCT CAGAATCAAG AAGACCTGCC CAGTTGGCCC CAGTTGGCCAC CTCCTCTGC CCAGATGAGA CTCCAGCCAT CTCCTTGCAGCAC CTCTTGAAGA CTGTTGAAGA CTGTTGAAGA	180 240 300 360 60 120 180 240 300 360 540 660 720 780 840 900 960 1020 1080 1140
45 50 55 60	TREAVVARLA LTELKLAIDG WVFNGRPHCF EGYVYYGLYL VRTDLNYIAA Nucleic Acid Acce Coding sequence: 1 GGAGCAGCCATA AGATGCACTT AAGACCCATA AGACCCATA AGAGCTTTCT TATGAATCCT TGGACATGTT AGAGGTTTAT AGAGTGCTC CAGATGCTC CAGATGCTC CAGTTATTT TCTCAGCAC TTCTAGAACAC TTCTAGAACAC TTCTAGAACAC TTCTAGAACAC TTCTAGAACAC TTCTAGAACAC GGTGAACCCT TCTATAAAAA GGTGAACCCT	DDMLEKLPPD TIIMSENLQD WMTGPFNPQG EGAGWDKRMM VDLRTAQTPE ssion #: AB033 68-3349 11 ACAACTTCAC GGAAGGAGAA CCTGAAAACA TCTGGAGCTG ACAACCCAAG TCTCATATCCAC CACAGACCT TATGCCAAGA GAAGTCTCCT TCAAAAAGTT TCCCAGGCTT TCCAAGACT CATGGTCACT CAAAAGTTACAC ACTCCTGAAG ATTATGAAAAGTT CCCAGGCTT CCAAGGCCATT AAAGTGCAAG ATTATGAAT AAAGTGCAAG ACTCCTGAAG ATTATGAAT AAAGTGGAGCCATT AAAGTGGAGCCATT AAAGTGGAGCCATT AAAGTGGAGCCATT	YVPFEVKERL ALDCMFDARI FLTAMRQEIT FLTAMRQEIT FLTEMRYVL E HWVLRGVA 036 Qunderlined sequ 21	QKMGPFQPMN PAWWKKASWV RANKGWALDN FELMPVIRTY LL CDVK SEC ences correspond 31	IFLRQEIDRM FSTLGFWFTE WVLCNEVTKW AENNTLRDPR ID NO:47 PDG5 to start and stop of the start and star	QRVLSLVRST LIERNSOFTS MKDDISTPPT FYSCPIYKKP DNA SEQUENCE dons) 51 TTTCAGATAA CATCAGCAGC CATACCATGA TGATGGTGGA CTCAGGTGTT CTACAGCAAA AAAAGCCTTC CTACAGCAAA AACAAGAAGT AGCTGGACCT CAGAATCAAG AACAACCAGC TCTCTCTCC CCAGTGGCCC TCTCCTCTCC	180 240 300 360 60 120 180 240 300 360 540 660 720 780 840 900 960 1020 1080 1140
45 50 55 60	TREAVVARLA LTELKLAIDG WVFNGRPHCF EGYVYVGLYL VRTDLNYIAA Nucleic Acid Acce Coding sequence: 1 GGAGCAGCCT AGATGACATG ACAGCCCATG AGACTCT TATGAATCT TATGAATCT TGATGCTGAA AGACTGCT CTCTCTCAGAA CAGATGCCT CAGTTATTT TCTCAGCAC TTCTATAATAT TCTCAGCCC TTCTATAAATAT TCTCAGCCC TTCTATAAACAC GGTGAACCCT GAGCATTTCT GAGCACC TTCTATAAACAC GTTAAAACAC GGTGAACCCT GAGCATTTCT GAGCACC TTCTATAAACAC GGTGAACCCT GAGCATTTCT GAGCACT TCTATAAACAC GGTGAACCCT GAGCATTTCT GAGCACT TCTATAAACAC GGTGAACCCT GAGCATTTCT GAGCACTTCT GAGCACTTCT GAGCACTTCT GAGCATTTCT GAGCACTTCT GAGCATTTCT GAGCACTTCT GAGCATTTCT GAGCATTTCT GAGCACTTTCT GAGCACTTTCT GAGCACTTTCT GAGCACTTTCT GAGCACTTTCT GAGCATTTCT GAGCACTTTCT GAGCACTTCT GAGCACTTTCT GAGCACTTTCT GAGCACTTTCT GAGCACTTTCT GAGCACTTTCT GAGCACTTTCT GAGCACTTCT GAGCACTTCT GAGCACTTCT GAGCACTTCT GAGCACTTCT GAGCACTTCT GAGCACTTCT GAGCACTTCT TCTCACTCT TCTCACTC TCTCACTCT TCTCACTC TCTCACTCT TCTCACTC TCTCC TCTCCT TCTCACTC TCTCC TCTC TCTCC TCTC T	DDMLEKLPPD TIIMSENLQD WMTGFFNPQG EGAGWDKRMM VDLRTAQTPE SSION #: AB033 68-3345 11 ACAACTTCAC GGAAGGAGAA TCTGGAGCTG ACAACCCAAG TCTCATTATCC CACGACCTT TCAAAAAAT TCCCAGGCTT TCAAAAAAT TCCCAGGCTT TCAAAAAAT TCCCAGGCTT TCAAAAAAT TCCCAGGCTT AAAATTATAAAT ACTGCTAAGA ACTCCTGAAG ACTCC	YVPFEVKERL ALDCMFDARI FLTAMRQEIT	QKMGPFQPMN PAWWKKASWV RANKGWALDN FELMPVIRTY LL CDVK SEC ences correspond 31	IFLRQEIDRM FSTLGFWFTE WVLCNEVTKW AENNTLRDPR ID NO:47 PDG5 to start and stop of CAGGGGGTTGC TCAGARARA GCATCACARA TCTCTCTCAC GCAGCAGAGG TTCAGCTTG ATTCAGCTTG GAGGAGGGGG GTTGAGGAGG TTTGAAGAGA CTCTCAGAGAG CAGCAGAGAG CAGCAGAGAG CAGCAGAGAG CAGCTGCCTT AGAGACCCTTG AGAGCCTTGA	QRVLSLVRST LIERNSQFTS MKXDDISTPPT FYSCPIYKKP DNA SEQUENCE odons) 51 TTTCAGATAA CATCAGCAGC CATACCATGA TGATGGTGGA CTCAGGTGTT ATTTACAAGA AAAAGCCTTC CTACAGCAA AACAAGCAGC ATGGTTCTGA AACAAGAAGT AGCTGGACCT CAGAATCAAG AAGACCTGCC CAGTTGGCCC CAGTTGGCCAC CTCCTCTGC CCAGATGAGA CTCCAGCCAT CTCCTTGCAGCAC CTCTTGAAGA CTGTTGAAGA CTGTTGAAGA	180 240 300 360 60 120 240 300 360 420 660 720 780 840 900 960 1020 1080 1140 1200
45 50 55 60 65 70	TREAVVARLA LTELKLAIDG WVFNGRPHCF EGYVYYGLYL VRTDLNYIAA Nucleic Acid Acce Coding sequence: 1 GGAGCAGCCT AGATGACATG ACAGCCCTATA AGATGCAGTC TATGAATCCT TGGAGATGTTCT TGGAGATGTTCT TGTTCAGAA CAGGTTATATA TTCTCAGCCC TTCTAAATAAT TTCTCAGCCC TTCTATAAAAA GGTGAACTCT GAGCATTTCT AGGCCTTCTATAAAAA GGTGAACTCCT TCTCAGCCC TTCTATAAAAA GGTGAACTCCT GAGCATTTCT AGGTGAACCACA	DDMLEKLPPD TIIMSENLQD WMTGFFNPQG EGAGWDERMM VDLRTAQTPE SSION #: AB033 68-3345 11 ACAACTTCAC GGAAGGAGAA CCTGAAAAC TCTGGAGCTG ACAACTCAGA CCTGAAAC TCTCATATCC AAAATGGAC TCACAGACCT ACATGTCACT ACAGGCGATT AAAGTGAAC ATTATGAATC CAGAGCGATT AAAGTGAGCCT AACATGTTCT AACATGTTCT AACATGTTCT AACATGTTCT AACATGTTCT AACATGTTCT AACATGTTCT AACATGTTCT	YVPFEVKERL ALDCMFDARI FLTAMRQEIT KLIESKPKVL HWVLRGVA 036 Qunderlined sequ 21 AACCAGAAAC ATGCTGGCAT AGGAGGCCAGA AGGAGGCCAGA AGTTAGAAGA CAGCTAGGAA TTACAGCAAG CTTCCCAGT TTGCTAGCA TTGCTTGCAGA CTTTCCCAGT CTTCCTAGT CTTGCTAGA CCTTTGGGAA CCTTTGGGAA CCTTTGGGAA CCTTTGGGAA CCTTTGGGAA CCTTTGGGAA CCTTTGGGAA CCTTTGGGAA CCTTTGGGAGC CAGAATTCA TGCTTGGAA CCTTTGGGAA CCTTTGGGAA CCTTTGGGAA CCTTTGGGAA CCTTTGGGAA CCTTTGGAA CCTGTTCA	QKMGPFQPMN PAWWKKASWV RANKGWALDN FELMPVIRTY LL CDVK SEC ences correspond 31	IFLRQEIDRM FSTLGFWFTE WVLCNEVTKW AENNTLRDPR ID NO:47 PDG5 to start and stop of 41 CAGGGGTTGC TCCAGAAAAG GATCACAAG GATCACAAG ATTTCAGTCTT GAGGAGGGGGG TTTGAAGAGT TTTGAAGAGT TCTGAAGAG GAGCAGAGAG GAGTCTCAA AGCATGAGAG CAGCTGCAT AGAGACACTT AGAGACCCTTG AGAGACAGAGAG CAGCCCTTGA GAGGCAGAGG CAGCCCTTGA	QRVLSLVRST LIERNSQFTS MKDDISTPPT FYSCPIYKKP DNA SEQUENCE odons) 51 TTTCAGATAA CATCAGCAGC CATACCATGA TGATGGTGGA CTCAGGTGTT ATTTACAAAA AAAAGCCTTC CTACAGCAAC CAAGGAAGCC ATGGTTCTGA AACAAGAAGT AGCTGGACCT CCAGATCACC CCAGATCCAC CCAGATCCCC CCAGTGGCAC TCCTCTCC CCAGTGGCAC TCCTCTTGCACAC TCCTCTTGCACAC TCCTCTTGACAC TCAGTTGGACAT TCATTTCTGT	180 240 300 360 120 120 240 300 360 420 720 780 840 900 1020 1080 1140 1200 1200
45 50 55 60 65 70	TREAVVARLA LTELKLAIDG WVFNGRPHCF EGYVYYGLYL VRTDLNYIAA Nucleic Acid Acce Coding sequence: 1 GGAGCAGCCT AGATGACATG AGATGCATG AGATGCATCT TATGAATCCT TGGAAATGT AGACTGTAA AGACTGCTT CAGTGCTT TCTCAGAA CAGATGCTT TCTCAGACAC TTCTAATAAT TTCTCAGCCC TTCTAATAAA GGTGAACCCT AGGCATTCT AGACACCC AGTTATTAAAA GGTGAACCCT AGTTCAACAA GGAGATTCT AGTTCAACAA GGAGATTCT AGTTCAACAA GGAGCCACTA	DDMLEKLPPD TIIMSENLQD WMTGFFNPQG EGAGWDKRMM VDLRTAQTPE SSION #: AB033 68-3345 11 ACAACTTCAC GGAAGGAGAA CCTGAAAACA TCTGGAGCTG ACAACCCAAGA CTTATATCC AAAATGGAGCT TATGCCAAGA GAGGTCTCT TCAAAAAGTT TCCAGGCTT TCCAGGCTT TCAAAAAGTT CAAAAAGTTACA ATTATGAATC CAGAGCGATT AAAGTGGAGC ATTATGAATC CAGAGCGATT AAAGTGGAGC ATGAACCT AAAATGTCT CAACATGTTCT AAAGTGGAGC ATGAACCT AACATGTTCT CACCCAGAT	YVPFEVKERL ALDCMFDARI FLTAMRQEIT KLIESKPKVL E HWVLRGVAI 036 0 (underlined sequ 21 AACCAGAAAA ATGCTGGCAT AGAAGACAGA AGAGACAGA AGAGACAGA AGAGACAGA CTTGCCAGT CTGCTGCAGA CTTGTTGAGGA CTTGTTGAGGA CTTGTTGAGGA CTTGTTGAGGA CTTGTTGAGGA CTTGTTGAGGA CTTGTTGAGGA CTGTTGAGGA CTGTTGGAAT CTGTTGAGGAAT CTGGGAAT CTGGGAAT CTGGGAAG CTGTGGAGCC AAGAGGTTCAGA CAGGTTCAGA CAGGTTCAGA CAGGTTCAGA CAGGTTCAGA CAGGTTCAGA CAGGTTCAGA CAGGTTCAGA	QKMGPFQPMN PAWWKKASWV RANKGWALDN FELMPVIRTY LL CDVK SEC ences correspond 31	IFLRQEIDRM FSTLGFWFTE WVLCNEVTKW AENNTLRDPR ID NO:47 PDG5 to start and stop of 1 CAGGGGTTGC TCCAGAAAG GATCCACAAC TCTCTCTCAC GCAGCAGAGG TTCAGCTTTG ATGACAGT TTTCAGTCTT CAGGGGGGG GAAGTCTTCA AGGCTCAGAGG CAACAAGAAG CAGCTCCTTTA AGACACCTT AGAGACCTTGA GGTTGAGAGG CAGCCCTTGA GGTTGAGAGG GATCCTTCAAA GGTTGAGAGG GATCCTTCAAA GGTTGAGAGG CAGCCCTTGA GGTTGAGAGG GATCCTTCAAA	QRVLSLVRST LIERNSOFTS MKDDISTPPT FYSCPIYKKP DNA SEQUENCE odons) 51 TITCAGATAA CATCAGCAGC CATACCATGA TGATGGTGGA CTCAGGTGTT CTACAGCAAA AAAAGCCTTC CTACAGCAAA AACAGAAGC AACCAGCAAA AACAGAAGC TCAGATCAAG AACAAGAAGC TCAGATCAAG AACAAGCAGC TCAGATCAAG TCAGATCAAG TCAGATCAAG TCAGATCAAG TCAGATCAAG TCAGATCAAG TCAGATCATC TCCAGCATG CTCTTGC CCAGTGGCAC TCCAGCCATG CTGTTGAAGA TCAATCCTAA TCACTCTGT TCCGGCAAAT	180 240 300 360 60 120 180 240 300 420 480 540 900 960 1020 1080 1140 1200 1320
45 50 55 60	TREAVVARLA LTELKLAIDG WVFNGRPHCF EGYVYYGLYL VRTDLNYIAA Nucleic Acid Acce Coding sequence: 1 GGAGCAGCCT AGATGACATO ACAGCCATA AGATGCAGCT TATGAATCCT TGGAAATGTT AGGAGATGTT AGGAGATGTT TCTCAGAA CAGTTATTT TCTCAGCAC TTCTAAAAA TCTCTAACAA GGTGAACCT GAGCATTCT AGTTCATCA AGTTCATCA TTCTAAAAA GGTGAACCCT TATAAAAA GGTGAACCCT TCTAAAAAA GGTGAACCCT TCTAAAAAA GGTGAACCCT GAGCATTTCT AGTTCAACAA GGTCACCACTA CTCAGAAAGC	DDMLEKLPPD TIIMSENLQD WMTGPFNPQG EGAGWDKRMM VDLRTAQTPE SSION #: AB033 68-3345 11 ACAACTTCAC GGAAGAGRA CCTGAAAACA TCTGGAGCTG ACAACCCAAG TCTCATATC CACAGACCT TATGCCAAGA CTGAAAAGTT TCCCAGGCTT TCCAGGCTT TCAAAAGTT TCCAGGCTT TCAAAAGTT TCCAGGCTT TCAAAAGTT TCCAGGCTT TAAAAGTT TCCAGGCTT CAAAAGTTATATATC ATGCTCAGG ATTATGAATC ATGAAGCCTC AAAATGTTATATATC CAGGCGATT AAAGTGGAGC ATGAAGCCTC AACATGTTCT TCCCCAGAT ACAGCTGTTG ACAGCTGTTG ACAGCTGTTG ACAGCTGTTG	YVPFEVKERL ALDCMFDARI FLTAMRQEIT KLIESKPKVL E HWVLRGVA 036 Qunderlined sequence AACCAGAAAC ATGCTGGCAA AGGAGGCCAI AGGAGGCCAI AGGAGGCCAI AGGAGGCCAI CAGCCAGGA CTTCCCAGCC CAGATTCAGA CTTCCCAGCC CAGATTCAGA CTTCTCGGAAA CACTTCTGA CACTTCTGA AGCAGAATGA CACTTCTGA CACTTCTGA AGCAGAATGA CACTTCTGA AGCAGATCAA AGCAGAATGA CACTTCTGA AGCAGATCAA AGCAGATCAA AGCAGATCAA AGCAGATCAA AGCAGATCAA AGCAGTTCAA ATCTCTCACA AGGAAGCCAC AGGAAGCCAC AGGAAGCCAC AGGAAGCCAC AGGAAGCAC AGGAAGCCAC AGGAAGCCAC AGGAAGCCAC AGGAAGCCAC AGGAAGCCAC AGGAAGCCAC AGGAAGCCAC AGGAAGCCAC AGGAAGCCAC	QKMGPFQPMN PAWWKKASWV RANKGWALDN FELMPVIRTY LL CDVK SEC ences correspond 31	IFLRQEIDRM FSTLGFWFTE WVLCNEVTKW AENNTLRDPR ID NO:47 PDG5 to start and stop of the start and star	QRVLSLVRST LIERNSQFTS MKDDISTPPT FYSCPIYKKP DNA SEQUENCE odons) 51 TTTCAGATAA CATCAGCAGC CATACCATGA TGATGGTGGA CTCAGGTGTT ATTTACAAAA AAAAGCCTTC CTACAGCAAC ATGGTTCTGA AACAAGAAGT AGCTGGCCATCAGGACCT CCAGATCAGC CCAGTGGCCC CCAGTGGCAC TCTCCTCTGC CCAGTTGGAAC TCATTCTGAAGA TCATTCTGA TCATTCTTA TCATTCTAA TCATTCTTAA TCATTCTTAC TCCAGCATAT CCAGATGCCT TCCAGCATAC TCCAGCATAC TCCAGCATAC TCCAGCATAT TCCAGCATAC TCCACATAC TCCACAC TC	180 240 300 360 60 120 180 240 300 360 420 480 540 600 720 780 900 960 1020 1020 1140 1200 1320 1380
45 50 55 60 65 70	TREAVVARLA LTELKLAIDG WVFNGRPHCF EGYVYVGLYL VRTDLNYIAA Nucleic Acid Acce Coding sequence: 1 GGAGCAGCCT AGATGACATG ACAGCCCATG AGACTGACATG AGACTGATC TATGAATCT TATGAATCT TGATGCTGAA AGACTGCT CAGTATGTT TCTCAGAA CAGTATGT TCTCAGAA GGTGACCT TCTATAATAT TCTCAGCCC TTCTATATAT TCTCAGCCC TTCTATATAT TCTCAGCAC TTCTATATAT TCTCAGCCC TTCTATATAT TCTCAGCAC TTCTATATAT TTCTCAGCAC TTCTATAAC GGTGAACCCT AGTTCACAC AGTTCACAC CAGGACACCT AGTTCACACA CTCAGAAAGC TTCCAGCACC TTCCAGCACC TTCCAGAAAGC TTCCCAGCCCC TTCCAGAAAGC TTCCCAGCCCC	DDMLEKLPPD TIIMSENLQD WMTGFFNPQG EGAGWDKRMM VDLRTAQTPE SSION #: AB033 68-3345 11 ACAACTTCAC GGAAGGAGAA CCTGAAAACA TCTGGACCTG ACAACTCACA GAAACTTCACT TCACAGACCT TATGCCAAGA CATGGTCACT TCAGACGTT TAAAATGTT TCCCAGGCTT AAAATGTAATC CAGACCGTT AAAATGTAATC CAGACGTTT AAAGTGGAGG ATGAAGCTT TCCCCAGAG ATGAAGCTT TCGGAGAGGGGC TCGGAGAGGGGGC TCGGAGAGGGGGC TCGGAGAGGGGGC TCGGAGAGGGGC	YVPFEVKERL ALDCMFDARI FLTAMRQEIT	QKMGPFQPMN PAWWKKASWV RANKGWALDN FELMPVIRTY LL CDVK SEC ences correspond 31	IFLRQEIDRM FSTLGFWFTE WVLCNEVTKW AENNTLRDPR ID NO:47 PDG5 to start and stop of CAGGGGTTGC TCAGARARA GCAGCAGAGG TTCAGCTTGA ATTCAGCTTG GAGGAGGGGG TTGAGGAGG TTGAGGAGG GAGCAGAGAG TTGAGGAGG GAGCAGAGAG CAGCAGAGAG CAGCAGAGAG CAGCAGAGAG CAGCTGCCTT AGAGACGC GAGCCTTGA GGTTGAGGAG GATCTCAGAG GGTTGAGGAG CAGCTGCCTT AGAGACCCTTGA GGTTGAGAGAG GATCCTCAAA CCGCTGCCTC AGTACTTCTG AGTACTTCTG AGTACTTCTG AGTACTTCTG AGTACTTCTG AGTACTTCTG AGTACTTCTG	QRVLSLVRST LIERNSQFTS MKDDISTPPT FYSCPIYKKP DNA SEQUENCE DOORS) 51 TTTCAGATAA CATCAGCAGC CATACCATGA TGATGGTGGA CTCAGGTGTT ATTTACABAA AAAAGCCTTC CTACAGCAGC CTACAGCAGC ATGGTTCTGA AACAGAAGT AGCTGGACCT CAGAATCAAG AAGACCTGC CCAGATGCCC CCAGTGGCCA TCTCCTCTCC	180 240 300 360 120 120 120 480 540 600 720 780 840 900 960 1020 1020 1140 1200 1320 1380 1440
45 50 55 60 65 70	TREAVVARLA LTELKLAIDG WVFNGRPHCF EGYVYYGLYL VRTDLNYIAA Nucleic Acid Acce Coding sequence: 1 GGAGCAGCCT AGATGCATA AGATGCATA AGATGCATA AGATGCATA AGATCCT TATGAATCCT TGGAATTCT TGATGCTTAAAAA CAGATGCTT TCTAAATAT TCTCAAGCCC TTCTAAAAA GGTGAACCT TTCTAAAAAA GGTGAACCT TCTAAAAAA GGTGAACCT TCTAAAAAA GGTGAACCT TCTAAAAAA GGTGAACCC TTCTAAAAAA CGTGAACCC TTCTAAAAAA CGTGAACCC TTCTAGAAACC CTCAGAAAGC CTCAGCCC CAGTCCTGTG	DDMLEKLPPD TIIMSENLQD WMTGFFNPQG EGAGWDKRMM VDLRTAQTPE SSION #: AB033 68-3345 11 ACAACTTCAC GGAAGGAGAA CCTGAAAACA TCTGGAGCTG ACAACCTAGAG TCTCATATCC AAAATGGAGT TATGCCAAGA CTTCAAAGAT TCCCAGAGCT TCAAAAGTT TCCCAGGT ACACTCTCAGAG ACTCTCAGAG ACTCCTCAGAG ACTCCTCAGAG ACTCCTCAGAG ACTCCTCAGAG ACTCCTCAGAG ACTCCTCAGAG ACTCCTCAGAG ACTCCTCAGAG ACTCCTCAGAG ACTACTCTCC CTCCCCAGAT ACAGCTCTTCC CTCCCCAGAT TCGCAGAGGGG GCACCAACAC	YVPFEVKERL ALDCMFDARI FLTAMRQEIT KLIESKPKVL E HWVLRGVA 36 (underlined sequ 21 AACCAGAAAC ATGCTGGCAI AGAAGACAGA AGAGAGCAGA CTTAGAGAA CTTGCCAGA CTTGCTAGA AGAGAGCAC CTTGCTGGGAAA AGCAGATCAGA CTTGTTGAGGA CTTGTTGAGGA CTTGTTGAGGA CTTGTTGAGGA CTTGTTGAGGA CTGCTCC CAGATTCTGA CTGCTCC AAGAAGATC CTAGGAAGA AGCAGAATGA CTTGCTAGA AGCAGATCAGA CTTCTGAGAGA CTTCTGAGAGA CTTCTGAGAGA CTTCTGAGAGA CTAGAGATCAGA CTAGAGATCAGA CTAGAGATCAGA CTAGAGATCCT CTAGAGATCCT CTAGAGATCCT CTAGAGATCCT CTAGAGATCCT CTTCCAAATF	QKMGPFQPMN PAWWKKASWV RANKGWALDN FELMPVIRTY LL CDVK SEC ences correspond 31 CACTACCCCT AGATTTCGGA CATGGTTAGT AGCAGAGCT AGCAGAGCT CAGAAGCT CATTCCCAG CACTTCATCCAC CACTTCTCCAC CACTTCTCAC CACTTCCCCG CAGCAATACAC CACTTCCCCCG CAGCAATACAC CACTTCCCCCG CAGCAATACAC CACTTCCCCCG CACCAATACAC CACTTCCCCCG CACCAATCCCCCCCCCC	IFLRQEIDRM FSTLGFWFTE WVLCNEVTKW AENNTLRDPR ID NO:47 PDG5 to start and stop of 41 CAGGGGTTGC TCCAGAAAG GATCCACAAC TCTCTCTCAC GCAGCAGTGG TTCAGCAGTG ATGACAAGT ATGACAAGT TCTGAGGAGG GAAGTTCA AGCTCAGAG GAAGATTCA AGCTCAGAG GAAGATTCA AGCTCAGAG GAAGATTCA AGCTCAGAG GAAGATTCAGAGG CAGCACTGCA AGACACTGC AGACACTTGA AGACACTTCTGA AGACACTTTGA AGACACTTCTGA AGACACTTCTGA AGACACTTCTGA AGACACTTTGA AGACACTTTGA AGACACTTTTGA AGACACTTTTGA AGACACTTTTGA AGACACTTTTTA	QRVLSLVRST LIERNSOFTS MKDDISTPPT FYSCPIYKKP DNA SEQUENCE dons) 51 TTTCAGATAA CATCAGCAGC CATACCATGA TGATGGTGGA TGATGGTGGA ACAAGAAGCTTC CTACAGCATC AACAAGAAGT AACAAGAAGT AACAAGAAGC TCAGATCAGC AACAAGAAGC TCAGATGCC TCTCCTCTGC CCAGTGGCCA TCAGATGCCT TCCAGCCATG TCAGATGCT TCCAGCCATG TCAGATGCT TCCAGCCATG TCAGATGCT TCCAGCATG CTGTTGAAGA TGAATCCTGT TCCAGCATG TCAGATGCCT TCAGATGCCT TCAGATGCCT TCAGATGCCT TCAGATGCAG TCAGATGCAT TCAGATGCAG CCCCTAAATT	180 240 300 360 120 120 240 300 360 420 720 780 840 900 1020 1080 1140 1200 1320 1320 1320 1340 1500
45 50 55 60 65 70	TREAVVARLA LTELKLAIDG WVFNGRPHCF EGYVYYGLYL VRTDLNYIAA Nucleic Acid Acce Coding sequence: 1 GGAGCAGCCCTATA AGATGCATGC AGATGCATCCT TATGAATCCT TGGAAATGTT AGAGCTGTA AGACTGCTCC CAGTTATATAT TTCTCAGCAC TTCTAGAAACC TTCTATAAAA GGTGAATCCT GAGCATTTCTAGCAC CTCTATAAAA GGTGAACCCT CTCTCAGAAACC TTCCAGCCC CTCTTATAAAA GGTGAACCCT CAGTTCACCT CAGCCCC CTCTCAGAAACC TTCCAGCCC CAGTCCTGTG TGAGAAACC TTCCAGCCC CAGTCCTGTG TGAGGAACTG	DDMLEKLPPD TIIMSENLQD WMTGPFNPQG EGAGWDKRNM VDLRTAQTPE SSION #: AB033 68-3345 11 ACAACTTCACA GGAAGGAGAA CCTGAAAACA TCTGAGACCCA ACAACCCAAG CTTCATATCCAG CACCAGACCT TATGCCAAGA TCTCAGACCT TCAAAAAGTT TCCCAGGCTT TCCAGGCTACT CAAAAAGTT TCCAGGCTACT CAAAAAGTT TCCAGGCTACT CAAAAAGTT CAAAAAGTT TCCAGGCTCAAG ATTATGAATC CAGAGCCATT AAAGTGGAGC ACACTGTTCC ACATGTTCC ACATGTTCC ACATGTTCC CTCCCCAGAT CTCCCCAGAT TCGGACAGCC GCACCAACAC TATCAACTCT TTCGGACAGCC TATCAACTCT TTCGGACAGCC TATCAACTCT TTCGGACAGCC TATCAACTCT TTCGGACACAC TATCAACTCT TTCGGACAGCC TATCAACTCT TTCGGACAGCC TATCAACTCT TTCGGACACCAC TATCAACTCT TTCGGACAGCC TATCAACTCT TTCGGACAGCC TATCAACTCT TTCAACTCT TTCGACAGCC TATCAACTCT TTCTCCCACACAC TTCTCCCACACAC TTCTCCCACACAC TTCTCCCACACAC TTCTCCCACACAC TTCTCCCACACAC TTCTCCCCACACAC TTCTCCCACACAC TTCTCCCCACACAC TTCTCCCCACACACA	YVPFEVKERL ALDCMFDARI FLTAMRQEIT KLIESKPKVL E HWVLRGVAJ 036 0 (underlined sequ 21 AACCAGAAAC ATGCTGGCAT AGAAGACAGA AGAGACAGA AGAGACAGA AGAGACAGA CTTCCCAGTC CTGCTCCC ACACTTCTGG CTTGGGAAA AGCAGAATCAGA CTTCTGGGAAA AGCAGAATCAGA CTTCTGGGAAA AGCAGAATCAGA CTTCTGGGAAA AGCAGAATCAGA CTTGGGAAA AGCAGAATCAG CTAGGGAGCC ACACTTCTGA CTTGGGAAA AGCAGAATCAG CTAGGGAGCC AGAGATCTC CAGGTTCAGA ATTCTCCTAGA ATTCTCCTAGA ATTCTCCTAGA CTGCACATCC CTGCAC	QKMGPFQPMN PAWWKKASWV RANKGWALDN FELMPVIRTY LL CDVK SEC ences correspond 31	IFLRQEIDRM FSTLGFWFTE WVLCNEVTKW AENNTLRDPR ID NO:47 PDG5 to start and stop of the start and start an	QRVLSLVRST LIERNSOFTS MKDDISTPPT FYSCPIYKKP DNA SEQUENCE DOORS) 51 TTTCAGATAA CATCAGCAGC CATACCATGA TGATGGTGGA TGATGGTGGA CTCAGGTGTT CAGAGCAAA AAAAGCCTTCC CAGAATCAAG AAGAACCAGC TCCTCTCTCC CAGGTGGCCC TCTCCTCTCC	180 240 300 360 60 120 180 240 300 360 420 480 540 960 1020 1080 1140 1260 1320 1380 1440 1560
45 50 55 60 65 70	TREAVVARLA LTELKLAIDG WVFNGRPHCF EGYVYYGLYL VRTDLNYIAA Nucleic Acid Acce Coding sequence: 1 GGAGCAGCCCTATA AGATGCATGC AGATGCATCCT TATGAATCCT TGGAAATGTT AGAGCTGTA AGACTGCTCC CAGTTATATAT TTCTCAGCAC TTCTAGAAACC TTCTATAAAA GGTGAATCCT GAGCATTTCTAGCAC CTCTATAAAA GGTGAACCCT CTCTCAGAAACC TTCCAGCCC CTCTTATAAAA GGTGAACCCT CAGTTCACCT CAGCCCC CTCTCAGAAACC TTCCAGCCC CAGTCCTGTG TGAGAAACC TTCCAGCCC CAGTCCTGTG TGAGGAACTG	DDMLEKLPPD TIIMSENLQD WMTGPFNPQG EGAGWDKRNM VDLRTAQTPE SSION #: AB033 68-3345 11 ACAACTTCACA GGAAGGAGAA CCTGAAAACA TCTGAGACCCA ACAACCCAAG CTTCATATCCAG CACCAGACCT TATGCCAAGA TCTCAGACCT TCAAAAAGTT TCCCAGGCTT TCCAGGCTACT CAAAAAGTT TCCAGGCTACT CAAAAAGTT TCCAGGCTACT CAAAAAGTT CAAAAAGTT TCCAGGCTCAAG ATTATGAATC CAGAGCCATT AAAGTGGAGC ACACTGTTCC ACATGTTCC ACATGTTCC ACATGTTCC CTCCCCAGAT CTCCCCAGAT TCGGACAGCC GCACCAACAC TATCAACTCT TTCGGACAGCC TATCAACTCT TTCGGACAGCC TATCAACTCT TTCGGACAGCC TATCAACTCT TTCGGACACAC TATCAACTCT TTCGGACAGCC TATCAACTCT TTCGGACAGCC TATCAACTCT TTCGGACACCAC TATCAACTCT TTCGGACAGCC TATCAACTCT TTCGGACAGCC TATCAACTCT TTCAACTCT TTCGACAGCC TATCAACTCT TTCTCCCACACAC TTCTCCCACACAC TTCTCCCACACAC TTCTCCCACACAC TTCTCCCACACAC TTCTCCCACACAC TTCTCCCCACACAC TTCTCCCACACAC TTCTCCCCACACAC TTCTCCCCACACACA	YVPFEVKERL ALDCMFDARI FLTAMRQEIT KLIESKPKVL E HWVLRGVAJ 036 0 (underlined sequ 21 AACCAGAAAC ATGCTGGCAT AGAAGACAGA AGAGACAGA AGAGACAGA AGAGACAGA CTTCCCAGTC CTGCTCCC ACACTTCTGG CTTGGGAAA AGCAGAATCAGA CTTCTGGGAAA AGCAGAATCAGA CTTCTGGGAAA AGCAGAATCAGA CTTCTGGGAAA AGCAGAATCAGA CTTGGGAAA AGCAGAATCAG CTAGGGAGCC ACACTTCTGA CTTGGGAAA AGCAGAATCAG CTAGGGAGCC AGAGATCTC CAGGTTCAGA ATTCTCCTAGA ATTCTCCTAGA ATTCTCCTAGA CTGCACATCC CTGCAC	QKMGPFQPMN PAWWKKASWV RANKGWALDN FELMPVIRTY LL CDVK SEC ences correspond 31	IFLRQEIDRM FSTLGFWFTE WVLCNEVTKW AENNTLRDPR ID NO:47 PDG5 to start and stop of the start and start an	QRVLSLVRST LIERNSOFTS MKDDISTPPT FYSCPIYKKP DNA SEQUENCE dons) 51 TTTCAGATAA CATCAGCAGC CATACCATGA TGATGGTGGA TGATGGTGGA ACAAGAAGCTTC CTACAGCATC AACAAGAAGT AACAAGAAGT AACAAGAAGC TCAGATCAGC AACAAGAAGC TCAGATGCC TCTCCTCTGC CCAGTGGCCA TCAGATGCCT TCCAGCCATG TCAGATGCT TCCAGCCATG TCAGATGCT TCCAGCCATG TCAGATGCT TCCAGCATG CTGTTGAAGA TGAATCCTGT TCCAGCATG TCAGATGCCT TCAGATGCCT TCAGATGCCT TCAGATGCCT TCAGATGCAG TCAGATGCAT TCAGATGCAG CCCCTAAATT	180 240 300 360 60 120 180 240 300 360 420 480 540 600 720 780 960 1020 1020 1140 1260 1320 1380 1440 1500
45 50 55 60 65 70	TREAVVARLA LTELKLAIDG WVFNGRPHCF EGYVYVGLYL VRTDLNYIAA Nucleic Acid Acce Coding sequence: 1 GGAGCAGCCT AGATGACATG ACAGCCATA AGATGCAGCT TATGAATCCT GGCCCAATCC GGCCCAATCC TCTCAGAA AGAACTGCT CAGATGCTT TCTCAGCAC TTCTAGAACAC TTCTAGAACAC TTCTAGACAC TTCTAGACAC TTCTAGACAC TTCTAGAACAC TTCTAACAAA GGTGAACCCT GAGCATTCT AGTTCACCC CAGCCCCCACTA CTCCAGCACCC CAGCCCCCCCCCC	DDMLEKLPPD TIIMSENLQD WMTGPFNPQG EGAGWDKRMM VDLRTAQTPF SSION #: AB033 68-3345 11 ACAACTTCAC GGAAGGAGAA CCTGAAAACA TCTGGAGCTG ACAACCCAAG TCTCATATCC CAAAATGGAGT TCATAGCCAGG CATGCTCAGG ACTCCTGAGG ACTGCTGAGG ACTGCTGAGG ACTGCTGAGG ACTGCTGAGG CTGCCAGAT CCTCCCAGAT ACACCTGTTG TCTCCCCAGAT ACACCTGTTG TCTCCCCAGAT ACACCTGTTG TCTCCCAGAT ACACCTGTTG TCTCCCCAGAT ACACCTGTTG TCTCCCCAGAT ACACCTGTTG TCTCCCCAGAT CTGCTTCCCAGT CTGCTCCAGT CTCCCAGT CTGCTCCAGT CTGCTCCAGT CTGCTCCAGT CTGCTCCAGT CTGCTCCAGT CTCCCAGT CTGCTCCAGT CTGCTCCAGT CTGCTCCAGT CTGCTCCAGT CTGCTCCAGT CTCCCAGT	YVPFEVKERL ALDCMFDARI FLTAMRQEIT KLIESKPKVL HWVLRGVA 36 Qunderlined sequence AACCAGAAAAC ATGCTGGCAA AGGAGGCCAI AGTTAGAAG CAGTTGCAGC CAGATTAGAG CTTCCCAGC CTTCCCAGC CTTCCTGGGAAA AGCAGAATGA CTTCCCAGC CTTCCCACC CTTCCACC CTCCCCCC CTCCCCCC CTCCCCCC CTCCCCCC CTCCCCCC	QKMGPFQPMN PAWWKKASWV RANKGWALDN FELMPVIRTY LL CDVK SEC ences correspond 31	IFLRQEIDRM FSTLGFWFTE WVLCNEVTKW AENNTLRDPR ID NO:47 PDG5 to start and stop of the start and st	QRVLSLVRST LIERNSOFTS MKDDISTPPT FYSCPIYKKP DNA SEQUENCE DOORS) 51 TTTCAGATAA CATCAGCAGC CATACCATGA TGATGGTGGA TGATGGTGGA CTCAGGTGTT CAGAGCAAA AAAAGCCTTCC CAGAATCAAG AAGAACCAGC TCCTCTCTCC CAGGTGGCCC TCTCCTCTCC	180 240 300 360 120 180 240 300 360 420 480 540 600 720 780 840 960 1020 1140 1200 1320 1380 1440 1500 1560 1620
45 50 55 60 65 70	TREAVVARLA LTELKLAIDG WVFNGRPHCF EGYVYVGLYL VRTDLNYIAA Nucleic Acid Acce Coding sequence: 1 GGAGCAGCCT AGATGACATG ACAGCCATA AGATGCAGCT TATGAATCCT GGCCCAATCC GGCCCAATCC TCTCAGAA AGAACTGCT CAGATGCTT TCTCAGCAC TTCTAGAACAC TTCTAGAACAC TTCTAGACAC TTCTAGACAC TTCTAGACAC TTCTAGAACAC TTCTAACAAA GGTGAACCCT GAGCATTCT AGTTCACCC CAGCCCCCACTA CTCCAGCACCC CAGCCCCCCCCCC	DDMLEKLPPD TIIMSENLQD WMTGPFNPQG EGAGWDKRMM VDLRTAQTPF SSION #: AB033 68-3345 11 ACAACTTCAC GGAAGGAGAA CCTGAAAACA TCTGGAGCTG ACAACCCAAG TCTCATATCC CAAAATGGAGT TCATAGCCAGG CATGCTCAGG ACTCCTGAGG ACTGCTGAGG ACTGCTGAGG ACTGCTGAGG ACTGCTGAGG CTGCCAGAT CCTCCCAGAT ACACCTGTTG TCTCCCCAGAT ACACCTGTTG TCTCCCCAGAT ACACCTGTTG TCTCCCAGAT ACACCTGTTG TCTCCCCAGAT ACACCTGTTG TCTCCCCAGAT ACACCTGTTG TCTCCCCAGAT CTGCTTCCCAGT CTGCTCCAGT CTCCCAGT CTGCTCCAGT CTGCTCCAGT CTGCTCCAGT CTGCTCCAGT CTGCTCCAGT CTCCCAGT CTGCTCCAGT CTGCTCCAGT CTGCTCCAGT CTGCTCCAGT CTGCTCCAGT CTCCCAGT	YVPFEVKERL ALDCMFDARI FLTAMRQEIT KLIESKPKVL HWVLRGVA 36 Qunderlined sequence AACCAGAAAAC ATGCTGGCAA AGGAGGCCAI AGTTAGAAG CAGTTGCAGC CAGATTAGAG CTTCCCAGC CTTCCCAGC CTTCCTGGGAAA AGCAGAATGA CTTCCCAGC CTTCCCACC CTTCCACC CTCCCCCC CTCCCCCC CTCCCCCC CTCCCCCC CTCCCCCC	QKMGPFQPMN PAWWKKASWV RANKGWALDN FELMPVIRTY LL CDVK SEC ences correspond 31	IFLRQEIDRM FSTLGFWFTE WVLCNEVTKW AENNTLRDPR ID NO:47 PDG5 to start and stop of the start and st	QRVLSLVRST LIERNSQFTS MKDDISTPPT FYSCPIYKKP DNA SEQUENCE odons) 51 TTTCAGATAA CATCAGCAGC CATACCATGA TGATGGTGGA CTCAGGTGTT ATTTACAAGA AAAAGCCTTC CTACAGCAA AAAAGCATTC CAGAATCAAG AGCACTGC ATGGTCTGA AGCAGCAGC CCAGATGACC CCAGATGACC CCAGATCAGC CCAGATCCC CCAGATGCCC CAGATGCCC CCAGATGCCC CCAGATTCCC CAGATTCCC CAGATTCCC CAGATTCCC CAGATTCCC CAGATTCCC CCAGATTCCC CCAGATTCCC CCCTAAATT AGGACATTTC AAGTCCAGCA GTCCATTGCC	180 240 300 360 120 180 240 300 360 420 480 540 660 720 780 900 960 1020 1080 1140 1200 1320 1380 1440 1500 1560 1620

	TCCCCAATAT	GCTACCCAGT	TCTTAAAGAG	GTCTAAAGTT	CAGGAAATGA	CCTCACGACT	1740
		GCTGTTGAAG					1800
	GTCATTCGTG	AAATTTATGG	CACAGCAAAT	CTTTTCAGAG	AGCTCTGCTC	TTAAGAGGGG	1860
						AGCCTGAAGT	1920
5	CAAGCACCAA	GTTTTCTCAG	ATTCAGGGAG	TGCTAATCCT	AAGGGAGGCA	TTTCTTCAAA	1980
						CACAGAAAGT	2040
						AGCTGTCTCC	2100
						CCCCTGTTTC	2160
10						AACATTCTTC	2220
10		GATAGGTCTA CAGAGCAGCG					2280 2340
		AGTTCTGTGA					2400
						TCAGCCAAGG	2460
						AGTTCACCAT	2520
15						AAGATCTTGA	2580
						ATGTTGAAAA	2640
						GTGAGAAACA	2700
	AGATAACTTC	ACCCAGCTTG	CTTCAGTGCC	CTCGGGCCCA	ATTTCATCCT	CTGTAGGCAG	2760
20						ACCTCACCAA	2820
20						TGGCCAAGAA	2880
						ATGCTGTCTC	2940
						CCCACATTTC	3000
						AGGAGCCTAA CTTCCAGTGT	3060 3120
25						CAGTTGGATT	3120
20		AAGATACTGC					3240
		AAGTTCCAGA					3300
						GAGCTCTTGT	3360
		AGCATTTÄTT					3420
30	CTCCCTCTCT	TACCCAGATT	GGAGTGCAGT	GGCGCGATCT	CCGCTCACTG	CAAGCTCCGC	3480
		CACGCCACTC					3540
						TTTCACCATG	3600
						CTCCCAAAAG	3660
35		AGGCGTGAGC				GTATTAAGCA	3720
JJ						TAAGCAAAAT	3780 3840
		ATGCAAATTA					3900
		GTAGTCACTG					3960
						TGTGCTTTAG	4020
40						CCCTCAACCA	4080
		TTCTAATGGA					4140
		AGATAGATGC					4200
		CGGTGGAAGT					4260
45						GCCCCTGATG	4320
43		GGGTGATAGT					4380
		GACCTGTCAC ATTATCTTAC					4440 4500
		TGTGACCCTC					4560
		AAGACCCTAT					4620
50		TGGCAAGCTT					4680
		CAGCATTAAG					4740
	TGTCTTTGCT	TCACAAGGTT	TTAGCCCAAA	GCAAGAGTGC	AATCCCAAAG	CCACAGAGAA	4800
	ATGAACTTTC	CGCTACCTGG	AAGCTTTAAG	TGAGTAAATC	AGCTTTTCCC	CTCTCATTCC	4860
e e						GTGACCCACT	4920
55						GAGGATGAGA	4980
						GCCTCCCTTC	5040
		ACACGGTCTT					5100 5160
	CACAMCCACA	ATGTAGGCAC CCACCCTAGA	AUCUCATOR	AGAGCCAGTT.	AACOURACTOR	ANTOTOTOTO	5220
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50 55 60	SEQ ID NO-50 P. Protein Accession 1 AFLSKVEEDD IEESKQETSM KIQTPELGEV AAAEPEDDSF MSSKLKSAQQ AAVLDDIQDL NVQVPEEPTH EPASVTPLEN IFLWRTVLVV NMILSDEAIK KDVISMIASE AELSEQIKSF	ABT Protein seque #: BAA13 11	21 NAINAKRSKE AAKGVNTGGR NDNPEEHLKT GHSDKREDLL LDKVFRASES AEETATLVMA ASEVSQKPNT YLTKSLVATL QISEKLKTIM NQBILDDTAK ARLSEEKVKS LTHKDDNINA	31 KNPGNQGRQF EPNTMVEKER SGLAGEPEGE ILSSFFKEQQ QILSIAEKML PPLEEGLGGA EKDLDPGPVT PDDVQPGPDF KENTELVQKL NLRVMLESER ECHRVQEENA LTNCITQLNL	DVNILQVPDRA PLADKKAQRP LSKEDHGNTE SLQRFQKYFN DTRVAENRDL MEEMQPLHED TEDTPMDAID YGLFWKPVPI SNYEQKIKES EQNVKNQDLI RLKKKKEQLQ LECESESEGQ	ULGTIHPDPE FERSDFSDSI KYMGTBSQGS VHELEALLQE GMNENNIFEE NFSREKTAEL ANKQPETAAE TAFLGIASFA KKHVQETRKQ SENKKSIEKL QEIEDWSKLH NKGGNDSDEL	120 180 240 300 360 420 480 540 600 660 720
50 55	SEQ ID NO:50 P. Protein Accession APLSKVEEDD IEESKQETSM KIQTPELGBV AAAEPEDDSF MSSKLKSAQQ AAVLDDIQDL NVQVPEEPTH IFLWRTVLVV NMILSDEAIK KDVISMIASE AELSEQIKSF ANGEVGGDRN	ABT Protein seque #: BAA13 11	21	31 KNPGNQGRQF EPNTMVEKER SGLAGEPEGE ILSSFFKEQQ QILSIAEKML PPLEEGLGGA EKDLDPGFVDT PDDVQPGPDF KENTELVQKL NLRVMLESER ECHRVQEEN LTNCITQLNL SVVEEDLKLL	DVNILQVPDRA PLADKKAQRP PLADKKAQRP LSKEDHGNTE SLQRFQKYFN DTRVAENRDL MEEMQPLHED TEDTPMDAID YGLPWKPVFI SNYEQKIKES EQNVKNQDLI RLKKKEQLQ LECESESEGQ QLKLRASVST	ULGTIHPDPE FERSDFSDSI KYMGTESQGS VHELEALLQE GMNENNIFEE MFSREKTAEL ANKQPETAAE TAFLGIASFA KKHVQETRKQ SENKKSIEKL QEIEDWSKLH NKGGNDSDEL KCNLEDQVKK	120 180 240 300 360 420 480 540 600 660 720 780
50 55 60	SEQ ID NO:50 P. Protein Accession APLSKVEEDD IEESKQETSM KIQTPELGEV AAAEPEDDST MSSKLKSAQQ AAVLDDIQDL NVQVPEEPTH EPASVTPLEN IFLWRTVLVV NMILSDEAIK KDVISMNASE AELSEQIKST ANGEVGGDRN LEDDRNSLQA	AB7 Protein seque #: BAA13 11 † ** PYSEELLEDE ILDSEKTSET FQNKDSDYLK HWTPHTSVE ESLPYNMEKV IYFVRYKHST LDQRVIGDTH AILLIYSFMF KDRVYQVTEQ YKDKIKTLEK FSEVQIALNE EKMKNQIKQM AKAGLEDECK	21 NAINAKRSKE AAKGVNTGGR NDNPEEHLKT GHSDKREDLL LDKVFRASES AEETATLVMA ASEVSOKPNT YLTKSLVATL QISEKLKTIM NQEILDDTAK ARLSEEKVKS LTHKDDNINA MDVSRTQTAI TLRQKVEILN	31 KNPGNQGRQF EFNTMVEKER SGLAGEPEGE GILSIAEKML PPLEEGLGGA EKDLDPGPVT PDDVQPGPPDF KENTELVQKL NLRVMLESER ECHRVQEENA LTNCITQLNL SVVEEDLKLL ELYQQKEMAL	DVNLQVPDRA PLADKKAQRP PLADKKAQRP LSKEDHGNTE SLQRFQKYFN DTRVAENRDL MEEMQPLHED TEDTPMDAID YGLPWKPVFI SNYEQKIKES EQNVKNQDLI RLKKKKEQLQ LECESESEGQ QLKLRASVST QKKLSQEEYE	ULGTIHPDPE FERSDFSDSI KYMGTESQGS VHELEALLQE GMNENNIFEE MFSREKTAEL ANKQPETAAE TAFIGIASPA KKHVQETRKQ SENKKSIEKL QEIEDWSKLH NKGGNDSDEL KCNLEDQVKK RQEREHRLSA	120 180 240 300 360 420 480 540 600 720 780 840
50 55 60	SEQ ID NO:50 P. Protein Accession AFLSKVEEDD IEESKQETSM KIQTPELGEV AAAEPEDDSF MSSKLKSAQQ AAVLIDDIQDL NVQVPEEPTH EPASVTPLEN IFLWRTVLVV NMILSDEAIK KDVISMNASE AELSEQIKSF ANGEVGGDRN LEDDRNSLQA ADEKAVSAAE	ABY Protein seque #: BAA13 11	21 MAINAKRSKE AAKGVNTGGR NDNPEEHLKT GHSDKREDLL LDKVFRASES AEETATLVMA ASEVSQKPNT YLTKSLVATL QISEKLKTIM NQEILDDTAK ARLSEEKVKS LTHKDDNINA MDVSRTQTAI TLRQKVEILN EMEDELQKTE	31 KNPGNQGRQF EPNTMVEKER SGLAGEPEGE IISSFFKEQQ QILSIAEKML PPLEEGLGGA EKDLDPGPVT PDDVQPGPDF KENTELVQKL NLRVMLESER ECHRVQEENA LTNCITQLNL SVVEEDLKLI RSFKNQIATH	DVNILQVPDRA PLADKKAQRP LSKEDHGNTE SLQRFQKYFN DTRVAENRDL MEEMQPLHED TEDTFMDAID YGLPWKPVFI SNYEQKIKES EQNVKNQDLI RLKKKKEQLQ LECESESEGQ QLKLRASVST QKKLSQEYE EKKAHENWLK	VLGTIHPDPE FERSDPSDSI KYMGTESGGS VHELEALLQE GMNENNIFEE NFSREKTAEL ANKQPETAAE TAPLGIASPA KKHVQETRKQ SENKKSIEKL QEIEDMSKLH NKGGNDSDEL KCNLEDQVKK RQEREHRLSA ARAAERAIAE	120 180 240 300 360 420 480 540 600 660 720 780
50 55 60	SEQ ID NO:50 P. Protein Accession AFLSKVEEDD IEESKQETSM KIQTPELGEV AAAEPEDDSF MSSKLKSAQQ AAVLIDDIQDL NVQVPEEPTH EPASVTPLEN IFLWRTVLVV NMILSDEAIK KDVISMNASE AELSEQIKSF ANGEVGGDRN LEDDRNSLQA ADEKAVSAAE	ABY Protein seque #: BAA13 11	21 NAINAKRSKE AAKGVNTGGR NDNPEEHLKT GHSDKREDLL LDKVFRASES AEETATLVMA ASEVSOKPNT YLTKSLVATL QISEKLKTIM NQEILDDTAK ARLSEEKVKS LTHKDDNINA MDVSRTQTAI TLRQKVEILN	31 KNPGNQGRQF EPNTMVEKER SGLAGEPEGE IISSFFKEQQ QILSIAEKML PPLEEGLGGA EKDLDPGPVT PDDVQPGPDF KENTELVQKL NLRVMLESER ECHRVQEENA LTNCITQLNL SVVEEDLKLI RSFKNQIATH	DVNILQVPDRA PLADKKAQRP LSKEDHGNTE SLQRFQKYFN DTRVAENRDL MEEMQPLHED TEDTFMDAID YGLPWKPVFI SNYEQKIKES EQNVKNQDLI RLKKKKEQLQ LECESESEGQ QLKLRASVST QKKLSQEYE EKKAHENWLK	VLGTIHPDPE FERSDPSDSI KYMGTESGGS VHELEALLQE GMNENNIFEE NFSREKTAEL ANKQPETAAE TAPLGIASPA KKHVQETRKQ SENKKSIEKL QEIEDMSKLH NKGGNDSDEL KCNLEDQVKK RQEREHRLSA ARAAERAIAE	120 180 240 300 360 420 480 540 600 720 780 840
50 55 60 65	SEQ ID NO-50 P. Protein Accession AFLSKVEEDD IEESKQETSM KIQTPELIEN MSSKLKSAQQ AAVLDDIQDL NVQVPEEPTH EPASVTPLEN IFLWRTVLVV NMILSDEAIK KDVISMNASE AELSEQIKSF ANGEVGGDRN LEDDRNSLQA ADERAVSAAE EKREAANLRH	ABT Protein seque #: BAA13 11	21 NAINAKRSKE AAKGVNTGGR NDNPEEHLKT GHSDKREDLL LDKVFRASES AEETATLVMA AEEVSQKPNT YLTKSLVATL QISEKLKTIM NQBILDDTAK AKLSEEKVKS LTHKDDNINA MDVSRTQTAI TLRQKVEILM MDVSRTQTAI TLRQKVEILM MLQEEPVIVK	31 KNPGNQGRQF EPNTMVEKER SGLAGEPEGE LISSFFKEQQ QILSIAEKML PPLEEGLGGA EKDLDPGPVT PDDVQPGPDF KENTELVQKL NLRVMLESER LTNCITQLNL SVVEEDLKLL ELYQQKEMAL ENSFKNQIATH PMPGKPNTQN	DVNILQVPDRA PLADKKAQRP LSKEDHGNTE SLQRFQKYFN DTRVAENRDL MEEMQPLHED TEDTPMDAID YGLFWKPVPI SNYEQKIKES EQNVKNQDLI RLKKKKEQLQ LECESESEGQ QLKLRASVST QKKLSQEEYE PRRGPLSQN	VLGTIHPDPE FERSDPSDSI KYMGTESGGS VHELEALLQE GMNENNIFEE NFSREKTAEL ANKQPETAAE TAPLGIASPA KKHVQETRKQ SENKKSIEKL QEIEDMSKLH NKGGNDSDEL KCNLEDQVKK RQEREHRLSA ARAAERAIAE	120 180 240 300 360 420 480 540 600 660 720 780 840 900
50 55 60 65	SEQ ID NO:50 P. Protein Accession APLSKVEEDD IEESKQETSM KIQTPELGEV AAAEPEDDSF MSSKLKSAQQ AAVLDDIQDL NVQVPEEPTH EPASVTPLEN IFLWRTVLVV NMILSDEAIK KDVISMNASE AELSEQIKSF ANGEVGGDRN LEDDRNSLQA ADEKAVSAAE EKREAANLRH GECSPPLTVE	ABT Protein seque #: BAA13 11 † YPSEELLEDE ILDSEKTSET FONKDSDYLK HWTPHTSVEP ESLPYNMEKV IYFVRYKHST LDQRVIGDTH AILLIYSFMF KDRVYQVTEQ YKDKIKTLEK FSEVQIALNE EKSQKDLEVA EKMKNQIKOM AKAGLEDECK EVKTYKRIIE EVKLELTQKMA PFVRPLSATL	21 NAINAKRSKE AAKGVNTGGR NDNPEEHLKT GHSDKREDLL LDKVFRASES AEETATLVMA ASEVSQKPNT YLTKSLVATL QISEKLKTIM NQEILDDTAK AKLSEEKVKS LTHKDDNINA MDVSRTQTAI TLRQKVEILN EMEDELQKTE MLQEEPVIVK NRRDMPRSEF	31 KNPGNQGRQF EPNTMVEKER SGLAGEPEGE ILSSFFKEQQ QILSIAEKML PPLEEGLGGA EKDLDPGPVT KENTELVQKL NLRVMLESER BCHRVQEENA LTNCITQLNL SVVEEDLKLL ELYQQKEMAL RSFKNQIATH PMPGKPNTQN GSVDGPLPHP	DVNIQVPDRA PLADKKAQRP PLADKKAQRP LSKEDHGNTE SLQRFQKYFN DTRVAENRDL MEEMQPLHED TEDTPMDAID YGLPWKPVFI SNYEQKIKES EQNVKNQDLI RLKKKEQLQ LECESESEGQ QLKLRASVST QKKLSQEYE EKKAHENWLK PPRRGPLSQN RWSAEASGKP	ULGTIHPDPE FERSDFSDSI KYMGTBSQGS VHELEALLQE GMNENNIFEE NFSREKTAEL ANKQPETAAE TAFLGIASFA KKHVQETRKQ SENKKSIEKL QEIEDWSKLH NKGGNDSDEL KCNLEDQVKK RQEREHRLSA ARAAERAIAE GSFGPSPVSG	120 180 240 300 360 420 480 540 600 660 720 780 840 900 960
50 55 60	SEQ ID NO:50 P. Protein Accession AFLSKVEEDD IEESKQETSM KIQTPELGEV AAAEPEDDST MSSKLKSAQQ AAVLDDIQDL NVQVPEEPTH EPASVTPLEN IFLWRTVLVV NMILSDEAIK KDVISMNASE AELSEQIKSF ANGEVGGDRN LEDDRNSLQA ADERAVSAAE EKREAANLRH GGCSPPLTVE TMMNSSSRGS	AB7 Protein seque #: BAA13 11 † ** PYSEELLEDE ILDSEKTSET FONKDSDYLK HWTPHTSVEP ESLPYNMEKV IYFVRYKHST LDQRVIGDTH AILLIYSFMF KORVYQVTEQ YKDKIKTLEK FSEVQIALNE EKMKNQIKOM AKAGLEDECK EVKTYKRRIE KLLELTQKME SPTRVLDEGK	21 NAINAKRSKE AAKGVNTGGR NDNPEEHLKT GHSDKREDLL LDKVFRASES AEETATLVMA ASEVSQKPNT YLTKSLVATL QISEKLKTIM NQEILDDTAK AKLSEEKVKS LTHKDDNINA MDVSRTQTAI TLRQKVEILN EMEDELQKTE MLQEEPVIVK NRRDMPRSEF	31 KNPGNQGRQF EFNTMVEKER SGLAGEPEGE IISSFFKEQQ QILSIAEKML PPLEEGLGGA EKDLDPGPVT PDDVQPGPPDF KENTELVQKL NLRVMLESER ECHRVQEENA LTNCITQLNL SVVEEDLKLL ELYQQKEMAL RSFKNQIATH PMPGKPNTQN GSVDGPLPHP PPGVPLMSTP	PLADKKAQRP PLADKKAQRP PLADKKAQRP LSKEDHGNTE SLQRFQKYFN DTRVAENRDI TEDTFMDAID TEDTFMDAID TEDTFMDAID TEDTFMDAID TELKKKEQLQ LECESESEQ QLKLRASVST QKKLSQEEYE EKKAHENWLK PPRRGPLSQN MWSAEASGKP MGGFVPPPIR	ULGTIHPDPE FERSDPSDSI KYMGTESQGS VHELEALLQE GMNENNIFEE MFSREKTAEL ANKQPETAAE TAFIGIASPA KKHVQETRKQ SENKKSIEKL QEIEDWSKLH NKGGNDSDEL KCNLEDQVKK RQEREIRLSA ARAAERAIAE GSFGPSFVSG SPSDPGSGTA YGPPPQLCGP	120 180 240 300 360 420 480 540 600 660 720 780 840 900 960 1020
50 55 60 65	SEQ ID NO:50 P. Protein Accession APLSKVEEDD IEESKQETSM KIQTPELGEV AAAEPEDDSF MSSKLKSAQQ AAVLDDIQDL NVQVPEEPTH EPASVTPLEN IFLWRTVLVV NMILSDEAIK KDVISMNASE AELSEQIKSF ANGEVGGDRN LEDDRNSLQA ADEKAVSAAE EKREANLRH GECSPPLTVE TMMNSSSRGS FGPRPLPPPF	ABY Protein seque #: BAA13 11 YPSEELLEDE ILDSEKTSET FONKDSDYLK HWTPHTSVEI ESLPYNMEKV IYFVRYKHST LDQRVIGDTH AILLIYSFMF KDRVYQVTEQ YKDKIKTLEK FSEVQIALNE EKSQKDLEVA EKMKNQIKQM AKAGLEDECK EVKTYKRRIE KLLELTQKMA PFVRPLSATL SPTRVLDEGK GPGMRPPLGL	21 MAINAKRSKE AAKGVNTGGR NDNPEEHLKT GHSDKREDLL LDKVFRASES AEETATLVMA ASEVSQKPNT YLTKSLVATL QISEKLKTIM NQEILDDTAK AKLSEEKVKS LTHKDDNINA MDVSRTQTAI TLRQKVEILN EMEDELQKTE MIQEEPVIVK NRRDMPRSEP VNMAPKGPPP REFAPGVPPG	31 KNPGNQGRQF EPNTMVEKER SGLAGEPEGE IISSFFKEQQ QILSIAEKML PPLEEGLGGA EKDLDPGPVT PDDVQPGPDF KENTELVQKL NLRVMLESER ECHRVQEENA LTNCITQLNIL ELYQKEMAL RSFKNQIATH PMPGKPNTQN GSVDGPLPHP PPGVPLMSTP RRDLPLHPRG	DVNILQVPDRA PLADKKAQRP LSKEDHGNTE SLQRFQKYFN DTRVAENRDL MEEMQPLHED TEDTFMDAID YGLPWKPVFI SNYEQKIKES EQNVKNQDLI RLKKKKEQLQ LECESESEGQ QLKLRASVSI QKKLSQEEYE EKKAHENWLK PPRRGPLSQN RWSAEASGK MGGFVPPPIR FLPGHAPFRP	VLGTIHPDPE FERSDPSDSI KYMGTESGGS VHELEALLQE GMNENNIFEE NFSREKTAEL ANKQPETAAE TAPLGIASPA KKHVQETRKQ SENKKSIEKL QEIEDMSKLH MKGGNDSDEL KCNLEDQVKK RQEREIRLSA ARAAERAIAE GSFGPSPVSG SPSDPGSGTA YGPPPQLCGP LGSLGPREYF	120 180 240 300 420 480 540 660 720 780 840 900 960 1020 1080
50 55 60 65	SEQ ID NO:50 P. Protein Accession APLSKVEEDD IEESKQETSM KIQTPELGEV AAAEPEDDSF MSSKLKSAQQ AAVLDDIQDL NVQVPEEPTH EPASVTPLEN IFLWRTVLVV NMILSDEAIK KDVISMNASE AELSEQIKSF ANGEVGGDRN LEDDRNSLQA ADEKAVSAAE EKREANLRH GECSPPLTVE TMMNSSSRGS FGPRPLPPPF	ABY Protein seque #: BAA13 11 YPSEELLEDE ILDSEKTSET FONKDSDYLK HWTPHTSVEI ESLPYNMEKV IYFVRYKHST LDQRVIGDTH AILLIYSFMF KDRVYQVTEQ YKDKIKTLEK FSEVQIALNE EKSQKDLEVA EKMKNQIKQM AKAGLEDECK EVKTYKRRIE KLLELTQKMA PFVRPLSATL SPTRVLDEGK GPGMRPPLGL	21 NAINAKRSKE AAKGVNTGGR NDNPEEHLKT GHSDKREDLL LDKVFRASES AEETATLVMA ASEVSOKPNT YLTKSLVATL QISEKLKTIM NQEILDDTAK ARLSEEKVKS LTHKDDNINA MDVSRTQTAI TLRQKVEILN EMEDELQKTE MLQEEPVIVE MRCMPRSEP VNMAPKGPPP	31 KNPGNQGRQF EPNTMVEKER SGLAGEPEGE IISSFFKEQQ QILSIAEKML PPLEEGLGGA EKDLDPGPVT PDDVQPGPDF KENTELVQKL NLRVMLESER ECHRVQEENA LTNCITQLNIL ELYQKEMAL RSFKNQIATH PMPGKPNTQN GSVDGPLPHP PPGVPLMSTP RRDLPLHPRG	DVNILQVPDRA PLADKKAQRP LSKEDHGNTE SLQRFQKYFN DTRVAENRDL MEEMQPLHED TEDTFMDAID YGLPWKPVFI SNYEQKIKES EQNVKNQDLI RLKKKKEQLQ LECESESEGQ QLKLRASVSI QKKLSQEEYE EKKAHENWLK PPRRGPLSQN RWSAEASGK MGGFVPPPIR FLPGHAPFRP	VLGTIHPDPE FERSDPSDSI KYMGTESGGS VHELEALLQE GMNENNIFEE NFSREKTAEL ANKQPETAAE TAPLGIASPA KKHVQETRKQ SENKKSIEKL QEIEDMSKLH MKGGNDSDEL KCNLEDQVKK RQEREIRLSA ARAAERAIAE GSFGPSPVSG SPSDPGSGTA YGPPPQLCGP LGSLGPREYF	120 180 240 300 420 480 540 660 720 780 840 900 960 1020 1080
50 55 60 65 70	SEQ ID NO:50 P. Protein Accession APLSKVEEDD IEESKQETSM KIQTPELGEV AAAEPEDDSF MSSKLKSAQQ AAVLDDIQDL NVQVPEEPTH EPASVTPLEN IFLWRTVLVV NMILSDEAIK KDVISMNASE AELSEQIKSF ANGEVGGDRN LEDDRNSLQA ADEKAVSAAE EKREANLRH GECSPPLTVE TMMNSSSRGS FGPRPLPPPF	ABY Protein seque #: BAA13 11 YPSEELLEDE ILDSEKTSET FONKDSDYLK HWTPHTSVEI ESLPYNMEKV IYFVRYKHST LDQRVIGDTH AILLIYSFMF KDRVYQVTEQ YKDKIKTLEK FSEVQIALNE EKSQKDLEVA EKMKNQIKQM AKAGLEDECK EVKTYKRRIE KLLELTQKMA PFVRPLSATL SPTRVLDEGK GPGMRPPLGL	21 MAINAKRSKE AAKGVNTGGR NDNPEEHLKT GHSDKREDLL LDKVFRASES AEETATLVMA ASEVSQKPNT YLTKSLVATL QISEKLKTIM NQEILDDTAK AKLSEEKVKS LTHKDDNINA MDVSRTQTAI TLRQKVEILN EMEDELQKTE MIQEEPVIVK NRRDMPRSEP VNMAPKGPPP REFAPGVPPG	31 KNPGNQGRQF EPNTMVEKER SGLAGEPEGE ILSSFFKEQQ QILSIAEKML PPLEEGLGGA EKDLDPGPVT KENTELVQKL NLRVMLESER BCHRVQEENA LTNCITQLNL SVVEEDLKLL ELYQQKEMAL RSFKNQIATH PMPGKPNTQN GSVDGPLPHP FPGVPLMSTP RRDLPLHPRG RDEPPPASQS	DVNILQVPDRA PLADKKAQRP PLADKKAQRP LSKEDHGNTE SLQRFQKYFN DTRVAENRDL MEEMQPLHED TEDTPMDAID YGLPWKPVFI SNYEQKIKES EQNVKNQDLI LECESESEGQ QLKLRASVST QKKLSQEEYE EKKAHENWLK PPRRGPLSQN RWSAEASGKP MGGFVPPPIR FLPGHAPFRP TSQDCSQALK	VLGTIHPDPE FERSDFSDSI KYMGTESQGS VHELEALLQE GMNENNIFEE MFSREKTAEL ANKQPETAAE TAPLGIASFA KKHVQETRKQ SENKKSIEKL QEIEDWSKLH MKGGNDSDEL KCNLEDQVKK RQEREIRLSA ARAAERAIAE GSFGPSFVSG SPSDPGSGTA YGPPPQLCGP LGSLGPREYF QSP	120 180 240 300 420 480 540 660 720 780 840 900 960 1020 1080
50 55 60 65 70	SEQ ID NO:50 P. Protein Accession AFLSKVEEDD IEESKQETSM KIQTPELGEV AAAEPEDDST MSSKLKSAQQ AAVLDDIQDL NVQVPEEPTH EPASVTPLEN IFLWRTVLVV NMILSDEAIK KDVISMNASE AELSEQIKST ANGEVGGDRN LEDDRNSLQA ADERAVSAAE EKREAANLRH GECSPPLTVE TMMSSSRGS FGPRPLPPPF IPGTRLPPPT	AB7 Protein seque #: BAA13 11 YPSEELLEDE ILDSEKTSET FQNKDSDYLK HWTPHTSVEP ESLPYNMEKV IYFVRYKHST LDQRVIGDTH AILLIYSFMF KDRUYQVTEQ YKDKIKTLEK FSEVQIALNE EKMKNQIKQM AKAGLEDECK EVKTYKRRIE KLLELTQKM SPTRVLDEGK GPGMRPPLGL HGPQEYPPPP	21 MAINAKRSKE AAKGVNTGGR NDNPEEHLKT GHSDKREDLL LDKVFRASES AEETATLVMA ASEVSQKPNT YLTKSLVATI NQEILDDTAK ARLSEEKVKS LTHKDDNINA MDVSRTQTAI TLRQKVEILN EMEDELQKTE MLQEEPVIVE MRDMPRSEP VNMAPKGPPP REFAPGVPSGS	31 KNPGNQGRQF EPNTMVEKER SGLAGEPEGE ILSSFFKEQQ QILSIAEKML PPLEEGLGGA EKDLDPGPVT KENTELVQKL NLRVMLESER BCHRVQEENA LTNCITQLNL SVVEEDLKLL ELYQQKEMAL RSFKNQIATH PMPGKPNTQN GSVDGPLPHP FPGVPLMSTP RRDLPLHPRG RDEPPPASQS	DVNILQVPDRA PLADKKAQRP PLADKKAQRP LSKEDHGNTE SLQRFQKYFN DTRVAENRDL MEEMQPLHED TEDTPMDAID YGLPWKPVFI SNYEQKIKES EQNVKNQDLI LECESESEGQ QLKLRASVST QKKLSQEEYE EKKAHENWLK PPRRGPLSQN RWSAEASGKP MGGFVPPPIR FLPGHAPFRP TSQDCSQALK	VLGTIHPDPE FERSDPSDSI KYMGTESGGS VHELEALLQE GMNENNIFEE NFSREKTAEL ANKQPETAAE TAPLGIASPA KKHVQETRKQ SENKKSIEKL QEIEDMSKLH MKGGNDSDEL KCNLEDQVKK RQEREIRLSA ARAAERAIAE GSFGPSPVSG SPSDPGSGTA YGPPPQLCGP LGSLGPREYF	120 180 240 300 420 480 540 660 720 780 840 900 960 1020 1080
50 55 60 65	SEQ ID NO-50 P. Protein Accession 1 AFLSKVEEDD IEESKQETSM KIQTPELGEV AAAEFEDDSF MSSKLKSAQQ AAVLDDIQDL INVQVPEEPTH EPASVTPLEN IFLWRTVLVV NMILSDEAIK KDVISMNASE AELSEQIKSF ANGEVGGDRN LEDDRNSLQA ADEKAVSAAE EKREAANLRH GECSPFLTVE TMMNSSRGS FGFRPLPPPF IPGTRLPPPT Nucleic Acid Acces	ABT Protein seque #: BAA13 11	21 NAINAKRSKE AAKGVNTGGR NDNPEEHLKT GHSDKREDLL LDKVFRASES AEETATLVMA ASEVSQKPNT YLTKSLVATL QISEKLKTIM NQBILDDTAK ARLSEEKVKS LTHKDDNINA MDVSRTQTAI TLRQKVEILM EMEDELQKTE MLQEEPVIVK NRRDMFRSEP VNMAPKGPPP REFAPGVPPG AVRDLLPSGS	31 KNPGNQGRQF EPNTMVEKER SGLAGEPEGE LISSFFKEQQ QILSIAEKML PPLEEGLGGA EKDLDPGPVT PDDVQPGPDF KENTELVQKL NLRVMLESER ECHRVQEENA LTNCITQLNL SVVEEDLKLL ELYQQKEMAL RSFKNQIATH PMPGKPNTQN GSVDGPLPHP PPGVPLMSTP RRDLPLHPRG RDEPPPASQS	DVNILQVPDRA PLADKKAQRP LSKEDHGNTE SLQRFQKYFN DTRVAENRDL MEEMQPLHED TEDTFMDAID YGLPWKPVFI SNYEQKIKES EQNVKNQDLI RLKKKKEQLQ LECESESEGQ QLKLRASVST QKKLSQEEYE EKKAHENWLK PPRGPLSQN RWSAEASGK MGGPVPPPIR FLPGHAPFRP TSQDCSQALK QID NO:51 PAB9	VLGTIHPDPE FERSDPSDSI KYMGTESGGS VHELEALLQE GMNENNIFEE NFSREKTAEL ANKQPETAAE TAPLGIASPA KKHVQETRKQ SENKKSIEKL QEIEDMSKLH NKGGNDSDEL KCNLEDQVKK RQEREHRLSA ARAAERAIAE GSFGPSPVSG SPSDPGSGTA VGSPEPQLCGP LGSLGPREYF QSP DNA SEQUENCE	120 180 240 300 420 480 540 660 720 780 840 900 960 1020 1080
50 55 60 65 70	SEQ ID NO:50 P. Protein Accession AFLSKVEEDD IEESKQETSM KIQTPELGEV AAAEPEDDST MSSKLKSAQQ AAVLDDIQDL NVQVPEEPTH EPASVTPLEN IFLWRTVLVV NMILSDEAIK KDVISMNASE AELSEQIKST ANGEVGGDRN LEDDRNSLQA ADERAVSAAE EKREAANLRH GECSPPLTVE TMMSSSRGS FGPRPLPPPF IPGTRLPPPT	ABT Protein seque #: BAA13 11	21 MAINAKRSKE AAKGVNTGGR NDNPEEHLKT GHSDKREDLL LDKVFRASES AEETATLVMA ASEVSQKPNT YLTKSLVATI NQEILDDTAK ARLSEEKVKS LTHKDDNINA MDVSRTQTAI TLRQKVEILN EMEDELQKTE MLQEEPVIVE MRDMPRSEP VNMAPKGPPP REFAPGVPSGS	31 KNPGNQGRQF EPNTMVEKER SGLAGEPEGE LISSFFKEQQ QILSIAEKML PPLEEGLGGA EKDLDPGPVT PDDVQPGPDF KENTELVQKL NLRVMLESER ECHRVQEENA LTNCITQLNL SVVEEDLKLL ELYQQKEMAL RSFKNQIATH PMPGKPNTQN GSVDGPLPHP PPGVPLMSTP RRDLPLHPRG RDEPPPASQS	DVNILQVPDRA PLADKKAQRP LSKEDHGNTE SLQRFQKYFN DTRVAENRDL MEEMQPLHED TEDTFMDAID YGLPWKPVFI SNYEQKIKES EQNVKNQDLI RLKKKKEQLQ LECESESEGQ QLKLRASVST QKKLSQEEYE EKKAHENWLK PPRGPLSQN RWSAEASGK MGGPVPPPIR FLPGHAPFRP TSQDCSQALK QID NO:51 PAB9	VLGTIHPDPE FERSDPSDSI KYMGTESGGS VHELEALLQE GMNENNIFEE NFSREKTAEL ANKQPETAAE TAPLGIASPA KKHVQETRKQ SENKKSIEKL QEIEDMSKLH NKGGNDSDEL KCNLEDQVKK RQEREHRLSA ARAAERAIAE GSFGPSPVSG SPSDPGSGTA VGSPEPQLCGP LGSLGPREYF QSP DNA SEQUENCE	120 180 240 300 420 480 540 660 720 780 840 900 960 1020 1080
50 55 60 65 70	SEQ ID NO:50 P. Protein Accession APLSKVEEDD IEESKQETSM KIQTPELGEV AAAEPEDDSP MSSKLKSAQQ AAVLDDIQDL NVQVPEEPTH EPASVTPLEN IFLWRTVLVV NMILSDEAIK KDVISMNASE AELSEQIKSF ANGEVGGDRN LEDDRNSLQA ADEKAVSAAE EKREAANLRH GECSPPLTVE TMMNSSSRGS FGPRPLPPPT IPGTRLPPPT Nucleic Acid Acce Coding sequence	ABT Protein seque #: BAA13 11	21 i NAINAKRSKE AAKGVNTGGR NDNPEEHLKT GHSDKREDLL LDKVFRASES AEETATLVMA ASEVSQKPNT YLTKSLVATL QISEKLKTIM NQEILDDTAK ARLSEEKVS LTHKDDINIA MIDVSRTQTAI TLRQKVEILM EMEDELQKTE MIQEEPVIVK NRRDMPRSEP VNMAPKGPPP REFAPGVPG AVRDLLPSGS 06457 4 (underlined sequ	31 KNPGNQGRQF EPNTMVEKER SGLAGEPEGE ILSSFFKEQQ QILSIAEKML PPLEEGLGGA EKDLDPGPVT PDDVQPGPDF KENTELVQKL NLRVMLESER BCHRVQEENA LTNCITQLNL SVVEEDLKLL ELYQQKEMAL RSFKNQIATH PMPGKPNTQN GSVDGPLPHP PRGVPLMSTP RRDLPLHPRG RDEPPPASQS SE ences correspond	DVNILOVPDRA PLADKKAQRP PLADKKAQRP PLADKKAQRP LSKEDHGNTE SLQRFQKYFN DTRVAENRDL MEEMQPLHED TEDTPMDAID YGLFWKPVFI SNYEQKIKES EQNVKNQDLI LECESESEGQ QLKLRASVST QKKLSQEEYE EKKAHENWLK PPRRGPLSQN RWSAEASGKP MGGFVPPPIR FLPGHAPFRP TSQDCSQALK Q ID NO:51 PAB9	VLGTIHPDPE FERSDFSDSI KYMGTESQGS VHELEALLQE GMNENNIFEE MFSREKTAEL ANKQPETAAE TAPLGIASFA KKHVQETRKQ SENKKSIEKL QEIEDWSKLH MKGGNDSDEL KCNLEDQVKK RQEREIRLISA ARAAERAIAE GSFGPSFVSG SPSDPGSGTA YGPPPQLCGP LGSLGPREYF QSP DNA SEQUENCE	120 180 240 300 420 480 540 660 720 780 840 900 960 1020 1080
50 55 60 65 70	SEQ ID NO-50 P. Protein Accession 1 AFLSKVEEDD IEESKQETSM KIQTPELGEV AAAEFEDDSF MSSKLKSAQQ AAVLDDIQDL INVQVPEEPTH EPASVTPLEN IFLWRTVLVV NMILSDEAIK KDVISMNASE AELSEQIKSF ANGEVGGDRN LEDDRNSLQA ADEKAVSAAE EKREAANLRH GECSPFLTVE TMMNSSRGS FGFRPLPPPF IPGTRLPPPT Nucleic Acid Acces	ABT Protein seque #: BAA13 11	21 NAINAKRSKE AAKGVNTGGR NDNPEEHLKT GHSDKREDLL LDKVFRASES AEETATLVMA ASEVSQKPNT YLTKSLVATL QISEKLKTIM NQBILDDTAK ARLSEEKVKS LTHKDDNINA MDVSRTQTAI TLRQKVEILM EMEDELQKTE MLQEEPVIVK NRRDMFRSEP VNMAPKGPPP REFAPGVPPG AVRDLLPSGS	31 KNPGNQGRQF EPNTMVEKER SGLAGEPEGE LISSFFKEQQ QILSIAEKML PPLEEGLGGA EKDLDPGPVT PDDVQPGPDF KENTELVQKL NLRVMLESER ECHRVQEENA LTNCITQLNL SVVEEDLKLL ELYQQKEMAL RSFKNQIATH PMPGKPNTQN GSVDGPLPHP PPGVPLMSTP RRDLPLHPRG RDEPPPASQS	DVNILQVPDRA PLADKKAQRP LSKEDHGNTE SLQRFQKYFN DTRVAENRDL MEEMQPLHED TEDTFMDAID YGLPWKPVFI SNYEQKIKES EQNVKNQDLI RLKKKKEQLQ LECESESEGQ QLKLRASVST QKKLSQEEYE EKKAHENWLK PPRGPLSQN RWSAEASGK MGGPVPPPIR FLPGHAPFRP TSQDCSQALK QID NO:51 PAB9	VLGTIHPDPE FERSDPSDSI KYMGTESGGS VHELEALLQE GMNENNIFEE NFSREKTAEL ANKQPETAAE TAPLGIASPA KKHVQETRKQ SENKKSIEKL QEIEDMSKLH NKGGNDSDEL KCNLEDQVKK RQEREHRLSA ARAAERAIAE GSFGPSPVSG SPSDPGSGTA VGSPEPQLCGP LGSLGPREYF QSP DNA SEQUENCE	120 180 240 300 420 480 540 660 720 780 840 900 960 1020 1080
50 55 60 65 70	SEQ ID NO-50 P. Protein Accession APLSKVEEDD IEESKQETSM KIQTPELGEV AAAEPEDDSF MSSKLKSAQQ AAVLDDIQDL NVQVPEEPTH EPASVTPLEN IFLWRTVLVV NMILSDEAIK KDVISMIASE AELSEQIKSF ANGEVGGDRN LEDDRNSLQA ADEKAVSAAE EKREAANLRH GECSPPLTVE TMMNSSSRGS FGPRPLPPPF IPGTRLPPPT Nucleic Acid Acce Coding sequence	ABT Protein seque #: BAA13 11	21 NAINAKRSKE AAKGVNTGGR NDNPEEHLKT GHSDKREDLL LDKVFRASES AEETATLVMA ASEVSQKPNT YLTKSLVATL QISEKLKTIM NQBILDDTAK AKLSEEKVKS LTHKDDNINA MDVSRTQTAI TLRQKVEILM MEMEDELQKTE MLQEEPVIVK NRRDMFRSEP VNMAPKGPPP REFAPGVPPG AVRDLLPSGS 06457 4 (undefined sequally	31 KNPGNQGRQF EPNTMVEKER SGLAGEPEGE LISSFFKEQQ QILSIAEKML PPLEEGLGGA EKDLDPGPVT PDDVQPGPDF KENTELVQKL MLRVMLESER LINCITQLNL SVVEEDLKLL ELYQQKEMAL RSFKNQIATH PMPGKPNTQN GSVIGPLPHP FPGVPLMSTP RRDLPJLHPRG RDEPPPASQS SE ences correspond	DVNILQVPDRA PLADKKAQRP LSKEDHGNTE SLQRFQKYFN DTRVAENRDL MEEMQPLHED TEDTFMDAID YGLPWKPVFI SNYEQKIKES EQNVKNQDLI RLKKKKEQLQ LECESESEGQ QLKLRASVST QKKLSQESYE EKKAHENWLK PPRGPLSQN RWSAEASGK MGGFVPPPIR FLPGHAPFRP TSQDCSQALK Q ID NO:51 PAB9 10 start and stop of	VLGTIHPDPE FERSDPSDSI KYMGTESGGS VHELEALLQE GMNENNIFEE NFSREKTAEL ANKQPETAAE TAPLGIASPA KKHVQETRKQ SENKKSIEKL QEIEDMSKLH NKGGNDSDEL KCNLEDQVKK RQEREHRLSA ARAAERAIAE GSFGPSPVSG SPSDPGSGTA YGPPPQLCGP LGSLGPREYF QSP DNA SEQUENCE	120 180 240 300 360 420 480 540 660 720 780 840 900 960 1020 1080 1140
50 55 60 65 70	SEQ ID NO-50 P. Protein Accession APLSKVEEDD IEESKQETSM KIQTPELGEV AAAEPEDDSF MSSKLKSAQQ AAVLDDIQDL NVQVPEEPTH EPASVTPLEN IFLWRTVLVV NMILSDEAIK KDVISMIASE AELSEQIKSF ANGEVGGDRN LEDDRNSLQA ADEKAVSAAE EKREAANLRH GECSPPLTVE TMMNSSSRGS FGPRPLPPPF IPGTRLPPPT Nucleic Acid Acce Coding sequence	ABT Protein seque #: BAA13 11	21 NAINAKRSKE AAKGVNTGGR NDNPEEHLKT GHSDKREDLL LDKVFRASES AEETATLVMA ASEVSQKPNT YLTKSLVATL QISEKLKTIM NQBILDDTAK AKLSEEKVKS LTHKDDNINA MDVSRTQTAI TLRQKVEILM MEMEDELQKTE MLQEEPVIVK NRRDMFRSEP VNMAPKGPPP REFAPGVPPG AVRDLLPSGS 06457 4 (undefined sequally	31 KNPGNQGRQF EPNTMVEKER SGLAGEPEGE LISSFFKEQQ QILSIAEKML PPLEEGLGGA EKDLDPGPVT PDDVQPGPDF KENTELVQKL MLRVMLESER LINCITQLNL SVVEEDLKLL ELYQQKEMAL RSFKNQIATH PMPGKPNTQN GSVIGPLPHP FPGVPLMSTP RRDLPJLHPRG RDEPPPASQS SE ences correspond	DVNILQVPDRA PLADKKAQRP LSKEDHGNTE SLQRFQKYFN DTRVAENRDL MEEMQPLHED TEDTFMDAID YGLPWKPVFI SNYEQKIKES EQNVKNQDLI RLKKKKEQLQ LECESESEGQ QLKLRASVST QKKLSQESYE EKKAHENWLK PPRGPLSQN RWSAEASGK MGGFVPPPIR FLPGHAPFRP TSQDCSQALK Q ID NO:51 PAB9 10 start and stop of	VLGTIHPDPE FERSDFSDSI KYMGTESQGS VHELEALLQE GMNENNIFEE MFSREKTAEL ANKQPETAAE TAPLGIASFA KKHVQETRKQ SENKKSIEKL QEIEDWSKLH MKGGNDSDEL KCNLEDQVKK RQEREIRLISA ARAAERAIAE GSFGPSFVSG SPSDPGSGTA YGPPPQLCGP LGSLGPREYF QSP DNA SEQUENCE	120 180 240 300 420 480 540 660 720 780 840 900 960 1020 1080

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                         1-864 (underlined sequences correspond to start and stop codons)
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20	SEQ ID NO:54 P Protein Accession	BH7 Protein sequ n #: FGENI	ence ESH predicted				
	1	11	21	31	41	51	
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20				SE	Q ID NO:55 PBJ5	DNA SEQUENCE	
30		ession #: AF381					
	Coding sequence	9: 33-137	(unaeninea seque	ences correspond	to start and stop co	odons)	
	1	11	21	31	41	51	
~~	Ī ·	Ī	Ī	1	Ī	Ī	
35				CGATGTGCTG			60
				AACTGAGAAA			120
				GAACTCAAGG TGAAAGTAAA			180 240
				AATCCTGTCA			300
40				TTTCTTATGT			360
						AACTCTCCAC	420
				TTTTCACGAT			480
	TAAGCCATAC	TTTATGTTCA	ATAAAAAGAG	AATAAGCAGG	A		
15							
45		BJ5 Protein seque					
	Protein Accession	n#: AAK8	3352				
	1 .	11	21	31	41	51	
	i .	î	i i	1	ĩ	i	
50	MCCETYYRLL	VLKMEKKSEE	LRNMDGLGNV	EKGH	•	•	
				SE	Q ID NO:57 PBJ7	DNA SEQUENCE	
		ession #: AA876					
55	Coding sequence	s: 1-2064	(undenined seque	ences correspond t	so start and stop or	odons)	
JJ	1	11	21	31	41	51	
	ī	ī	ī	ī	ī	Ī	
	ATGGACAGTT	GCCTGCAACA	TATGAGAGAC	CTACTTTACC	TCCTTCAGGA	GCTCAGGTGT	60
60	TTAAATCCAG	CTACACTACT	CCCTGATCCA	GACTCCACTA	CTCCTGTTCA	TGACTGTCAG	120
60				CCTGATCTTC			180
				AGCTTCCTCG			240
				CAGGAAGGAC		AGAAGAAAAA AAACACTTACT	300 360
				TTTGCAGTTG			420
65				AATTTGCCGG			480
				AGCCAAACTG			540
						CCCTGACGCT.	
				CCTGATTGGA			660
70				CTTTCCATAA ACTATAACTG			720
70				AGACTTTATA			780 840
				GTCTCATGGA			900
				CAGAAACACC			960
						TCAACCCAGC	
<i>75</i>						TAAACTAGCC	1080
	CAAGATTGTT	GGCTATGTTT	AAAAGCAAAA	CCCCCTTATT	ATGTAGGATT	AGGAGTAGAA	
						CACAATAGGA	1200
						TGCTTCTCCT	
80						CTCTTACCAA	1320
		WITH THE STREET	THE RESERVE TO SERVE	TURGUTUTUR	CICGCIGCAT	TAATGGAACT	1380

			CGTGTTAGTT				1440
			CATCGCTCCC				1500
			ATTGGCTGGT AGAAACTGGA				1560 1620
5			TGCCATAGAT				1680
			CCGATGCTTA				1740
			TTGTTGCTTC				1800
			AAATCTAGAT TAACTGGAAC				1860 1920
10			ACTATTAAGT				1980
			ACGCATAGCT				2040
	TATGACACCC	TTGTTAATAA	C <u>TGA</u>				
	SEO IO NO EO D	BJ7 Protein seque		•			
15	Protein Accession		ESH predicted				
	1	11	21	31	41	51	
	MDSCLOHMRD	LLYLLOELRC	 LNPATLLPDP	DSTTPVHDCO	DLLETTKTGO	PDLODVPLEK	60
20	ADATVFTDGS	SFLEQGERKA	VSFPQPDLPD	NPTYSTEEEK	LASDVGANKN	QEGRVFANTT	120
			EPARTHEEQH				180
			SCRDTYQFFC				240 300
			QWYYGMSWGL VPLPFLVPRP				360
25			ATLKRGPLSC				420
			APNNTWLACT				480
			VPLLVPLLAG EVVLQNCRCL				540 600
			PWYOSMFNWN				660
30		SVKLTYLKTQ	_				•••
				or.	0.10.110.50.000	DVA OFOUTNOT	
	Nucleic Acid Acce	ession#: NM_0	19005	35	יטאר פב:טא עורט	I DNA SEQUENCE	•
0.5	Coding sequence		85 (underlined sec	quences correspon	d to start and stop	codons)	
35	_						
	1	11	21	31	41	51 1	
	TGATGGTGGA	AATTTCTTGA	AACCGCTCTC	GTAATTTGCC	ACGTGCTGTT	GCAAATATTC	60
40			AGCATGGCTT				120
40			GTGAATGGAC				180
			ATATTTTATG TTTATCATGT				240 300
			AAGACTCTGC				360
45			CCTGGTATCT				420
43			TTGTACTTAC AGTTTGTTCC				480 540
	CTGGAATCCA	CTGGATAGTA	ACTGGCTAGC	TGCTGGTTTA	GATAAGCACA	GAGCTGACTT	600
	TTCAGTGCTA	ATATGGGATA	TCTGCAGCAA	ATATACTCCT	GATATAGTTC	CCATGGAAAA	660
50			AAACTGAAAC				720
30			GTCTGTCTCT ACCTAGCTAT				780 840
	GTTCGTAAAT	ACAAAAGCTG	TTCAGGGTGT	GACGGTAGAC	CCATATTTCC	ACGATCGTGT	900
			AGGTTGCAAT				960
55			CAAAACCCTT				1020 1080
-			ATGAAACTGA				1140
			CCTTTGCGTG				1200
			TGTCAGACTT TGTGGGCTTG				1260 1320
60			AAAAAGATAT				1380
	AAGGTATGGA	CTTGATACAG	AGCAGGTGTG	GAGGAACCAC	ATTTTAGCTG	Gaaatgaaga	1440
			GGTATACTCT				1500
			ACAAAGGATC TGGAAAGCAG				1560 1620
65			ATGAAGAGAG				1680
			ACGTGGGGCC				1740
			TGGCATTGTT CTGAAAAAGG				1800 1860
			GATGAGAAGA				1920
70	TGCGATTACA	GCTAAATAAC	CCGTATTTGT	GTGTCATGTT	TGCATTTCTG	ACAAGTGAAA	1980
			TTGTATGAAA				2040
			GATACTCAGA				2100
			AATTTTGCTT AACTGGAGAT				2160 2220
75	GGTTCACCTT	TAGATGTTCT	TAAAGATGAA	AGGGTTCAGT	ACTGGATTGA	GAATTATAGA	2280
			GTTTTGGCAT				2340
			GCCTTTAGCA TTCAGCTGTG				2400 2460
			GAAATCTAAA				2520
80			TCTCATTAAT				2580

5	AACTGGTTTA TGGTTCAGGG GATACAACGG AGAGAACCCT	CATGGTGTCA ACCATGCAGA GGAATCTGGT TCAAGTGTGG CCATTCATGA	TAATTGCAGG GTGCCCTGTG ACCTGCAGAG AGCTTTCTAG	CACGGTGGAC TCTGCATGCA ACTGTCCAGC TAGGTGTCCT	AAAAATTAGC ATGCTGGACA CGTGTAAATG CATAAAATGT TCATAGCTCA AAATCATTCT	TATGCTTAGT TATGCAGTTG TACCACCTTA GAAACATACC	2640 2700 2760 2820 2880 2940
10	SEQ ID NO:60 Protein Accession	CQ1 Protein sequents: NP_06					
	1	11	21	31	41	51	
15	PYMKCVAWYL WNPLDSNWLA LGQNDACLSL ASFYEGQVAI	NYDPECLLAV AGLDKHRADF CWLPRDQKLL WDLRKFEKPV	GQANGRVVLT SVLIWDICSK LAGMHRNLAI LTLTEQPKPL	SLGQDHNSKF YTPDIVPMEK FDLRNTSQKM TKVAWCPTRT	GSLRLSEDSA KDLIGKEFVP VKLSAGETET FVNTKAVQGV GLLATLTRDS	KHARQCNTLA TLLVTKPLYE TVDPYFHDRV NIIRLYDMQH	60 120 180 • 240 300
20	SPITSLMWAC POLKSLWYTL	GRHLYECTEE HFMKQYTEDM ILALQLCGWI	ENDNSLEKDI DOKSPGNKGS KKGTDVDVGP	ATKMRLRALS LVYAGIKSIV	VTPNRTMSDF RYGLDTEQVW KSSLGMVESS WERAAAVALF	RNHILAGNED RHNWSGLDKQ	360 420 480 540
25			_	SE	Q ID NO:61 PDG3	DNA SEQUENCE	
	Nucleic Add Acce Coding sequence			ences correspond	to start and stop o	odons)	
30	1	11	21	31	41	51	
	GATCAGCCCA GAGTCCTGGC	CAGTACACAT	CATTGATGAC GACTTATAAA	AATTTCACTO	GTCTCAACCT	ATGGTGTGAT TTCTCATGCT ATTAAGAGAT TAAAGAATGT	120 180
35	CCATATTGTT TGTGATTTGG CTGAGCCTCA	GTATTTCATT ACCATGGCAC GTTTTCCTCA	GTGGTGTATT TTAAAAACTC TTTTCAAAT	GGAAAGTGAT TATAACCTCA ATAGAGAGTA	CTGGACTTTG GGCAAGTCTT TAACATTTAT	ATTATTTTT AGTGAGAAGA TTAATCTTCT CTCATAAGAC AGATTCCATA	360 420 480
40	TTCCAGTCTT TACCAAAAGA CAAATATACT AAATATTTTT	CTGACACGTO TTCTTTAACT	GTTTATCTGC GAGTTAATCA TCTGTGTGGG TTAGGAAAAA	CACAGGGAGA TTTGACAGAT TATCACTTAG TTGTAGNCAA	AGTCCTCAGA GCAAATGCTT GGAAAAAAAG ATCTTTTINI	TAAAAATGTC CCACCCCAA GCAGGCAACA CCCATTAACA AAACAGGTAA	600 660 720 780
45	CTTACTTGA		TCAMGGGGTA	. AIAAAAAIA	. AAAGICIICC	MANCAGGIAA	840
	SEQ ID NO:62 P						
50	1	11	21	31	41	51	
		DKFRKFIKAP	PRNYSMIVMF	TALQPQRQCS	VCRQANEEYQ	ILANSWRYSS	60 120
55	WIADRTDVHI AMTSGQMWNH	RVFRPPNYSG IRGPPYAHKN	TIALALLVSL PHNGQVSYIH	VGGLLYXRRN	KRADTFDLOR NLEFIYNKTG SHIILVLNAA PYSDLDFE	WAMVSLCIVE	180 240 300
60	Nucleic Acid Acce	Perion # AT NRO	235	SEC	2 ID NO:63 PDG8	DNA SEQUENCE	
00	Coding sequence				to start and stop c	odons) -	
	1	11 	21 	31 	41 	51 	
65	GGGGCGCCCA CAGGGCGAGC CCGGGCTGGC	CCGCGCTGCCCCCCCCCCCCCCCCCCCCCCCCCCCCCC	AGCCTACCCC CTGCTGCCTA TAAGCCCATT	GCGGCCGAGC GACTTCAGCC GAGTCCACGC	CGCCCGGGCC TGGAGGAGCT TGGTGGCCTG	CGCCACCGCC GCTGTGGCTG GCAGGGCGAG CTTCATGACC CATCGCCGGC	180 240
70	TTCCTGCCCA GCCGCAGTGC GCGGCCGTCA CTGTGTCCGC	ACGGCATGGA CCGCAGGGAC CTTCGGGGGT GCGCGCGGGT	ACAGCGCCGG CACCGCAGCC GGCGACCAAG GCCTTTCCCG	ACCACCGCCA GCCGCCGCCG TGACCCGCTC CCGGGGACTC	GCACCACCGC CCGCCGCTGC CGCTCCTCCC GGCCGGTGTG	AGCCACCCC CGCCGCCGCC TGTGTCCGTC CTTCGTGCTG	360 420 480 540
75	AAGCAAGGTT GGGTGGGATC CATCCCCTG CCAAATCCTG	TGTGCTGCGC GCAGCTCCGA CCCGACTCCT TGTATTGTT	TTCCAGTTCC AGACGGAGAG TCCCCGCACC TTATATATTT	GAAAAGCAGA GAGGGAAATG CACGTGCCCT AATAACTGTT	GGGCCCTTTC AGATTCATGG TTAAATGAAA	CGCCTTTGAA CTTGGACTGA CCCTCTATTG CAGAAAATGA GTTTTAGTAA	660 720 780 840
80						CTCTTTGTAT CGGGGGAGGG	

	TCCAAGTTG		GACGTGGCC	r GGTGGGCGT	r TCTTCTTGT/	T ATGAACCTTT A CTTATGTGGT	
5	SEQ ID NO:64 P Protein Accession	DG8 Protein seque					
	1	11	21	31	41	51	
10	PGWRLNRKPI	 AAPSAAAATA ESTLVACFMT AAAAAAAA	LVIVVWSVAA	LIWPVPIIAG		 DFSLEELQGE TTASTTAATP	60 120
15	Musicia Acid Accus	ession#: NM_0	00708	SE	Q ID NO:65 PDM1	DNA SEQUENCE	
13	Coding sequence		95 (underlined sec	quences correspon	d to start and stop	codons)	
oo .	1	11	21 .	31	41	51	
20	TCCCGGAGGC	CCGGGTCCCT TGGCCGGGCA	GGCGTGGTGC	GCGGTAGGAG	CTGGGCGCGC	ACGGCTACCG	60 120
		AGACACTGCC CGGCGGCTGC					180 240
25	GCTGCTGCTC	TGCATCCAGC	TCGGGGGAGG	ACAGAAGAAA	AAGGAGAATC	TTTTAGCTGA	300
		CAGCTGATGG AAATTTATAA					360 420
	TGCTCTTCAG	CCTCAGCGGC	AGTGTTCTGT	GTGCAGGCAA	GCTAATGAAG	AATATCAAAT	480
30		TCCTGGCGCT GAGGGGACAG					540 600
	CAYGCATTTW	CCTCCAAAAG	GCAGACCTAA	GAGAGCTGAT	ACTITIGACC	TCCAAAGAAT	660
		GCTGAGCAACT					720 780
25	TGGAGGTTTG	CTTTATTNGA	GAAGGAACAA	CTTGGAGTTC	ATCTATAACA	AGACTGGTTG	840
35		TCTCTGTGTA CCATATGCTC					900 960
						ATGCCGCTAT	1020
						TTGGAAAAAG	1080 1140
40						TTCTACTTTC AGTGAGAAGA	1200
	TGTGATTTGG	ACCATGGCAC	TTAAAAACTC	TATAACCTCA	GCTTTTTAAT	TAAATGAAGC	1260
						TGACTTTATA ACTTTTTTTA	1320 1380
15	AACTGTGGGT	TTTCCTAGTA	AATTTAATTT	ACAGAAATCA	ATGGTAGCAT	TTAGTAATCT	1440
45		TATCAAAGTG CAATGTAATT		CTGTTATATY	CAGTGTGTKC	CACAGGATTG	1500
50	SEQ ID NO:66 P Protein Accession	DM1 Protein sequents: NP_00		•			
	1	11	21	31 	41 	51 	
55						EKVEQLMEWS	60
23		DKFRKFIKAP VDYDEGTDVF					120 180
	WIADRTDVHI	RVFRPPNYSG	TIALALLVSL	VGGLLYXRRN	NLEFIYNKTG	WAMVSLCIVE	240
		IRGPPYAHKN RRIICLVGLG				ITMGMVLLNE	300
60	AATSKGDVGK	idizzen de de	BVVIIIBIBB	JII KOKINGI	11000010		
				QE.	∩ IN NA-67 POM	DNA SEQUENCE	
		ession#: NM_0					
65	Coding sequence	: 88-161	7 (underlined sequ	iences correspond	to start and stop o	codons)	
05							
	1 . I	11	21	31	41	51 I	
7 0		GAACTCTCCC			TGCCACCGTT		60
70		TCTTGCAGGT					120
						TTACTTGCAG TAGAGTTAAA	180 240
	TIGITAAAAT	CAGTTGAAAA	TCTTGGAGTG	AGCTATGTGA	AAGGAACTGA	ACAATACCAG	300
75						GCTAGAAGAT TGCTTATTGC	360 420
, ,						TCGATTTAGA	
	TTTAGTATTT	TACCCAAGGA	TAAAATTCAG	GATTTCTTAA	AGGATAGCCA	ATTGCAGTTT	540
						TGCTGATGCT	600 660
80						CGTACCACTT	720
						_	

	AACCACATTC	TOCOLADOCAT	ССТСАВТСАВ	TTTAGAGCCA	ממשששמה	CCCTTTTCCCA	780
				TCTGATGAAA			840
				TACAGTACCC			900
_				AAATCCTTCC			960
5	CATAAAGCCT	TGCGGGAAAA	TCACCATCTT	CGTCATGGAG	GCCGAATGCA	GTATGGCCTA	1020
	TTTCTGAAGG	GCATTGGTTT	AACTTTGGAA	CAGGCATTGC	AGTTCTGGAA	GCAAGAATTT	1080
				GATAAAGGTT			1140
				TATACACCTT			
							1200
10				CATGGGTGCC			1260
10	GAGCTGCTGA	AGCAAAAGTT	GCAGTCATAC	AAGATCTCTC	CTGGAGGGAT	AAGCCAGATT	1320
	TTGGATTTAG	TAAAGGGGAC	ACATTACCAG	GTAGCCTGTC	AAAAATACTT	TGAGATGATA	1380
	CACAATGTGG	ATGATTGTGG	CTTTTCTTTG	AATCATCCTA	ATCAGTTCTT	TTGTGAGAGC	1440
				AAGAAGGAAC			1500
				GATGCATCAT			1560
15							
13				GAAGATTACT			1620
	GTTTTATAAC	CCTTTTTCCT	CANTAGCCTG	TTTCCTGTTT	TTAAGATTTT	CCTTTGTTG	1680
	TTGAAAAAGG	GTTTCACTGT	CACCAAGGCT	TAGTGCAGTG	ACACAATTAC	AGCTGATTGC	1740
	AGCCTTGACC	TTCCCAGCTC	AAGTGATCCT	CCTACCTCAG	CCTCCCAAGT	ACTTACCACA	1800
				TTTTTCAATT			1860
20							
20				ACTCCTGGGC			1920
				GAGCCACTGT			1980
	TAACCTTTTC	GTTTAACTTC	TCTCTTCACT	GCATCCCAAT	CCATCTACAG	GCATGCACAC	2040
	TTATTAGGAA	AGGAGGTTTG	AGGTAACAAC	AGAGACTTTC	ACTATATTTT	GCTTTGACAG	2100
				TGTCACTTGA			2160
25				GGTTAAAGTC			2220
				TGTAATTTTG			
				TGTAATTTTG	ACTCAATCCT	TITCIGGACC	2280
	ATTTTTGTTA	ATAAATATCA	AAGTGT				
30	SEQ ID NO:68 PI	M2 Protein senue	nce:	•			
-	Protein Accession						
	LIORGII MOSSIOII	W. NF_00	0936	•			
	_						
	1	11	21	31	41	51	
~ ~	1	1			i		
35	MEFSGRKRRK	LRLAGDORNA	SYPHCLOPYL	OPPSENISLT	EPENLAIDRV	KLLKSVENLG	60
				DEYEPRRRDH			120
				FEAISDEEKT			180
							240
				LKDIVAIILN			
	QSDERLQPLL	NHLSHSYTGO	DYSTOGNVGK	ISLOOTDILLS	TKSFPPCMRO	THKALRENHH	300
40							
40		LFLKGIGLTL	EQALQFWKQE	FIKGKMDPDK	FDKGYSYNIR	HSPGKEGKRT	360
40	LRHGGRMQYG DYTPFSCLKI	LFLKGIGLTL	EQALQFWKQE	FIKGKMDPDK	FDKGYSYNIR	HSPGKEGKRT	
40	DYTPFSCLKI	LFLKGIGLTL ILSNFPSQGD	EQALQFWKQE YHGCPFRHSD	FIKGKMDPDK PELLKQKLQS	FDKGYSYNIR YKISPGGISQ	HSFGKEGKRT ILDLVKGTHY	360
40	DYTPFSCLKI QVACQKYFEM	LFLKGIGLTL ILSNPPSQGD IHNVDDCGFS	eqalqfwkqe Yhgcpfrhsd Lnhpnqffce	FIKGKMDPDK	FDKGYSYNIR YKISPGGISQ	HSFGKEGKRT ILDLVKGTHY	360 420
40	DYTPFSCLKI	LFLKGIGLTL ILSNPPSQGD IHNVDDCGFS	eqalqfwkqe Yhgcpfrhsd Lnhpnqffce	FIKGKMDPDK PELLKQKLQS	FDKGYSYNIR YKISPGGISQ	HSFGKEGKRT ILDLVKGTHY	360 420
	DYTPFSCLKI QVACQKYFEM	LFLKGIGLTL ILSNPPSQGD IHNVDDCGFS	eqalqfwkqe Yhgcpfrhsd Lnhpnqffce	FIKGKMDPDK PELLKQKLQS SQRILNGGKD	FDKGYSYNIR YKISPGGISQ IKKEPIQPET	HSPGKEGKRT ILDLVKGTHY PQPKPSVQKT	360 420
45	DYTPFSCLKI QVACQKYFEM KDASSALASL	LFLKGIGLTL ILSNFPSQGD IHNVDDCGFS NSSLEMDMEG	EQALQFWKQE YHGCPFRHSD LNHPNQFFCE LEDYFSEDS	FIKGKMDPDK PELLKQKLQS SQRILNGGKD	FDKGYSYNIR YKISPGGISQ IKKEPIQPET	HSFGKEGKRT ILDLVKGTHY	360 420
	DYTPFSCLKI QVACQKYFEM KDASSALASL Nucleic Acid Acce	LFLKGIGLTL ILSNFPSQGD IHNVDDCGFS NSSLEMDMEG . ssion#: NM_02	EQALQFWKQE YHGCPFRHSD LNHPNQFFCE LEDYFSEDS 24840	FIKGKMDPDK PELLKQKLQS SQRILNGGKD SEC	FDRGYSYNIR YKISPGGISQ IKKEPIQPET 1 ID NO:69 PDM3	HSFGKEGKRT ILDLVKGTHY PQPKPSVQKT DNA SEQUENCE	360 420
	DYTPFSCLKI QVACQKYFEM KDASSALASL	LFLKGIGLTL ILSNFPSQGD IHNVDDCGFS NSSLEMDMEG . ssion#: NM_02	EQALQFWKQE YHGCPFRHSD LNHPNQFFCE LEDYFSEDS 24840	FIKGKMDPDK PELLKQKLQS SQRILNGGKD	FDRGYSYNIR YKISPGGISQ IKKEPIQPET 1 ID NO:69 PDM3	HSFGKEGKRT ILDLVKGTHY PQPKPSVQKT DNA SEQUENCE	360 420
	DYTPFSCLKI QVACQKYFEM KDASSALASL Nucleic Acid Acce Coding sequences	LFLKGIGLTL ILSNPPSQGD IHNVDDCGFS NSSLEMDMEG ssion #: NM_07 108-491	EQALQFWKQE YHGCPFRHSD LANHPNQFFCE LEDYFSEDS 4840 (underlined sequ	FIKGKMDPDK PELLKQKLQS SQRILNGGKD SEX ences correspond	FDKGYSYNIR YKISPGGISQ IKKEPIQPET 2 ID NO:69 PDM3 to start and slop c	HSPGKEGKRT ILDLVKGTHY PQPKPSVQKT DNA SEQUENCE odons)	360 420
45	DYTPFSCLKI QVACQKYFEM KDASSALASL Nucleic Acid Acce	LFLKGIGLTL ILSNFPSQGD IHNVDDCGFS NSSLEMDMEG . ssion#: NM_02	EQALQFWKQE YHGCPFRHSD LNHPNQFFCE LEDYFSEDS 24840	FIKGKMDPDK PELLKQKLQS SQRILNGGKD SEC	FDRGYSYNIR YKISPGGISQ IKKEPIQPET 1 ID NO:69 PDM3	HSFGKEGKRT ILDLVKGTHY PQPKPSVQKT DNA SEQUENCE	360 420
	DYTPFSCLKI QVACQKYFEM KDASSALASL Nucleic Acid Acce Coding sequences	LFLKGIGLTL ILSNPPSQGD IHNVDDCGFS NSSLEMDMEG ssion #: NM_07 108-491	EQALQFWKQE YHGCPFRHSD LANHPNQFFCE LEDYFSEDS 4840 (underlined sequ	FIKGKMDPDK PELLKQKLQS SQRILNGGKD SEX ences correspond	FDKGYSYNIR YKISPGGISQ IKKEPIQPET 2 ID NO:69 PDM3 to start and slop c	HSPGKEGKRT ILDLVKGTHY PQPKPSVQKT DNA SEQUENCE odons)	360 420
45	DYTPFSCIKI QVACQKYFEM KDASSALASL Nucleic Acid Acce Coding sequence:	LFLKGIGLTL ILSNPFSQGD IHNVDDCGFS NSSLEHDMEG ssion #: NM_07 108-491	EQALQFWKQE YHGCPFRHSD LINHPNQFFCE LEDYFSEDS 4840 (underlined sequ	FIKGKMDPDK PELLKQKLQS SQRILNGGKD SEX ences correspond	FDKGYSYNIR YKISPGGISQ IKKEPIQPET ID NO:69 PDM3 to start and slop c 41	HSPGKEGKRT ILDLVKGTHY POPKPSVQKT DNA SEQUENCE odons) 51	360 420
45	DYTPFSCIKI QVACQKYFEM KDASSALASL Nucleic Acid Acce Coding sequence: 1 AATTCATACA	LFLKGIGLTL ILSNPPSGGD IHNVDDCGFS NSSLEHDMEG SSION #: NM_02 108-491 11	EQALQFWKQE YHGCPFRHSD LNHPNQFFCE LEDYFSEDS 4840 (underlined sequ 21 CATATATATATG	FIKGKMDPDK PELLKQKLQS SQRILNGGKD SEC ences correspond 31	FDKGYSYNIR YKISPGGISQ IKKEPIQPET 2 ID NO:69 PDM3 to start and stop c 41 GGAAAAGGCT	HSPGKEGKRT ILDLVKGTHY PQPKPSVQKT DNA SEQUENCE cdons) 51 TCATCAAGAA	360 420 480
45	DYTPFSCIKI CVACQKYFEM KDASSALASL Nucleic Acid Acce Coding sequence: 1 AATTCATACA GTCTCGGCTC	LFLKGIGLTL ILSNPPSQGD IHNVDDCGFS NSSLEMDMEG SSION#: NM_07 108-491 11	EQALQFWKQE YHGCPFRHSD LNHPNQFFCE LEDYFSEDS A840 (underlined sequ 21 CATATATATAG AGAGAGTTCA	FIKGKMDPDK FELLKQKLQS SQRILNGGKD SEC ences correspond 31	FDKGYSYNIR YKISPGGISQ IKKEPIQPET 2 ID NO:69 PDM3 to start and stop c 41 GGARAAGGCT AAACCACATG	HSPGKEGKRT ILDLVKGTHY PQPKPSVQKT DNA SEQUENCE odons) 51 TCATCAAGAA GATGCAGCCT	360 420 480 60 120
45	DYTPFSCIKI CVACQKYFEM KDASSALASL Nucleic Acid Acce Coding sequence: 1 AATTCATACA GTCTCGGCTC GTGTGGGAAG	LFLKGIGLTL ILSNPFSQGD IHNVDDCGFS NSSLEHDMEG ssion #: NM_07 108-491 11 GGAGAGAAGT ATTAATCATC GGCCTTCTCCA	EQALQFWKQE YHGCPFRHSD LNHPNQFFCE LEDYFSEDS 4840 (underlined sequ 21 CATATATATAG AGAGAGTTCAG AAAGGTTCAG	FIKGKMDPDK PELLKQKLQS SQRILNGGKD SEX ences correspond 31 CAGTGATTGT TACAGGAGAG GCTCACTGAA	FDKGYSYNIR YKISPGGISQ IKKEPIQPET DID NO:69 PDM3 to start and stop c 41 GGAAAAGGCT AAACCACATG CACCACAGAA	HSPGKEGKRT ILDLVKGTHY PQPKPSVQKT DNA SEQUENCE odons) 51 TCATCAAGAA GATGCAGCCT CTCATACAGG	360 420 480 60 120 180
45	DYTPFSCIKI CVACQKYFEM KDASSALASL Nucleic Acid Acce Coding sequence: 1 AATTCATACA GTCTCGGCTA AGGGAAG	LFLKGIGLTL ILSNPPSGGS NSSLEMDMEG SSION #: NM_0Z 108-491 11 GGAGAGAAGT ATTAATCATC GGCTTCTCCA TATGAATGCA	EQALQFWKQE YHGCPFRHSD LINHPNQFCE LEDYFSEDS 4840 (underlined sequ 21 CATATATATA AGAGATCAA AAAGGTCCAA CTGAATGTGA	FIKGKMDPDK PELLKQKLQS SQRILNGGKD SEC ences correspond 31 CAGTGATTGT TACAGGAGAG GCTCACTGAA CAAAGCATTC	FDKGYSYNIR YKISPGGISQ IKKEPIQPET ID NO:69 PDM3 to start and stop c 41 GGAAAAGGCT AACCACAGAA CGCTGGAAAA	HSPGKEGKRT ILDLVKGTHY PQPKPSVQKT DNA SEQUENCE odons) 51 TCATCAAGAA GATGCAGCCT CTCATACAGG CACAGCTCAA	360 420 480 60 120 180 240
45	DYTPFSCIKI CVACQKYFEM KDASSALASL Nucleic Acid Acce Coding sequence: 1 AATTCATACA GTCTCGGCTC GTGTGGGAAG AGAAGAACCA TGCACATCAG	LFLKGIGLTL ILSNPPSGED IHNVDDCGFS NSSLEHDMEG SSION #: NM_02 108-491 11 GGAGAGAGAGT GGAGAGAGGT ATTAATCATC GCCTTCTCCA AAAGCTCACA	EQALQFWKQE YHGCPFRHSD LNHPNQFFCE LEDYFSEDS 4840 (underlined sequ 21 CATATATATAT AGAGAGTTCA AAAGGTTCAG CTGAATAGAA	FIKGKMDPDK FELLKQKLQS SQRILNGGKD SEC ences correspond 31	FDKGYSYNIR YKISPGGISQ IKKEPIQPET 2 ID NO:69 PDM3 to start and stop c 41 GGAAAAGGCT AAACCACATG CACCAGAAA CGCTGGAAAT	HSPGKEGKRT ILDLVKGTHY PQPKPSVQKT DNA SEQUENCE odons) 51 TCATCAAGAA GATGCAGCTC CTCATACAGG CACAGCTCAA GTGGAAAAGG	360 420 480 60 120 180 240 300
45	DYTPFSCIKI CVACQKYFEM KDASSALASL Nucleic Acid Acce Coding sequence: 1	LFLKGIGLTL ILSNPPSQGD IHNVDDCGFS NSSLEMDMEG SSION #: NM_02 108-491 11	EQALQFWKQE YHGCPFRHSD LNHPNQFFCE LEDYFSEDS A840 (underlined sequ 21 CATATATATA AGAGATTCA AAAGGTCCAG CAGAGAGAA TCATTGTACA	FIKGKMDPDK FELLKQKLQS SQRILNGGKD SEC ences correspond 31	FDKGYSYNIR YKISPGGISQ IKKEPIQPET 2 ID NO:69 PDM3 to start and stop c 41 GGARAAGGCT AAACCACATG CACCAGGAA CGCTGGAAAT TGCCGTGAATT CATACTGGAG	HSPGKEGKRT ILDUVKGTHY PQPKPSVQKT DNA SEQUENCE odons) 51 TCATCAAGAA GATGCAGCCTA CTCATACAGG CACAGCCTCAA GTGGAAAAGG AAAACCCTA	360 420 480 60 120 180 240 300 360
45	DYTPFSCIKI CVACQKYFEM KDASSALASL Nucleic Acid Acce Coding sequence: 1 AATTCATACA GTCTCGGCTC GTGTGGGAAG AGAGAAGCCC TCCACATCAG CTTCATTCAG TATATGCAAT	LFLKGIGLTL ILSNPFSQGD IHNVDDCGFS NSSLEHDMEG SSION #: NM_07 108-491 11 GGAGAGAAGT ATTAATCATC TATGAATGCA AAAGCTCACA AAAGCTACAC GAATGTGGAA	EQALQFWKQE YHGCPFRHSD LNHPNQFFCE LEDYFSEDS 4840 (underlined sequ 21 CATATATATG AGAGAGTTCA AAAGGTCCAG CTGAATGTAA TCATTGTACA AAGGCTTCAT	FIKGKMDPDK PELLKQKLQS SQRILNGGKD SEC ences correspond 31 CAGTGATTGT TACAGGAGAG GCTCACTGAA CAAAGCATTC GTCATATATA TCAGCGAATT CCAAAAGGGC	FDKGYSYNIR YKISPGGISQ IKKEPIQPET ID NO:69 PDM3 to start and stop c 41 GGAAAAGGCT AAACCACATG CACCAGAGAA CGCTGGAAAT TGCCGTGATC CATACTGGAG AACCTCCTTA	HSPGKEGKRT ILDLVKGTHY PQPKPSVQKT DNA SEQUENCE odons) 51 TCATCAAGAA GATGCAGCCT CTCATACAGG CACAGCTCAA GTGGAAAAG GTGGAAAAG TTCATCGACG TTCATCGACG	360 420 480 60 120 180 240 300 420
45	DYTPFSCIKI CVACQKYFEM KDASSALASL Nucleic Acid Acce Coding sequence: 1 AATTCATACA GTCTCGGCTC GTGTGGGAAG AGAGAAGCCC TCCACATCAG CTTCATTCAG TATATGCAAT	LFLKGIGLTL ILSNPFSQGD IHNVDDCGFS NSSLEHDMEG SSION #: NM_07 108-491 11 GGAGAGAAGT ATTAATCATC TATGAATGCA AAAGCTCACA AAAGCTACAC GAATGTGGAA	EQALQFWKQE YHGCPFRHSD LNHPNQFFCE LEDYFSEDS 4840 (underlined sequ 21 CATATATATG AGAGAGTTCA AAAGGTCCAG CTGAATGTAA TCATTGTACA AAGGCTTCAT	FIKGKMDPDK FELLKQKLQS SQRILNGGKD SEC ences correspond 31	FDKGYSYNIR YKISPGGISQ IKKEPIQPET ID NO:69 PDM3 to start and stop c 41 GGAAAAGGCT AAACCACATG CACCAGAGAA CGCTGGAAAT TGCCGTGATC CATACTGGAG AACCTCCTTA	HSPGKEGKRT ILDLVKGTHY PQPKPSVQKT DNA SEQUENCE odons) 51 TCATCAAGAA GATGCAGCCT CTCATACAGG CACAGCTCAA GTGGAAAAG GTGGAAAAG TTCATCGACG TTCATCGACG	360 420 480 60 120 180 240 300 360
45 50 55	DYTPFSCIKI CVACQKYFEM KDASSALASL Nucleic Acid Acce Coding sequence: 1 AATTCATACA GTCTCGGCTC GTGTGGGAAG AGAGAAGCCC TGCACATCAG CTTCATTCAG TATATGCAAT TACTCACACT	LFLKGIGLTL ILSNPFSGGS INNUDCGFS NSSLEHDMEG SSION #: NM_02 108-491 11	EQALQFWKQE YHGCPFRHSD LNHPNQFFCE LEDYFSEDS 4840 (underlined sequ 21 CATATATATA AGAGATTCA AAAGGTCCAG CTGAATGTGA CAGGAGAGAA TCATTGTACA AAGGCTTCAT CCTATGAATG	FIKGKMDPDK FELLKQKLQS SQRILNGGKD SEC ences correspond 31	FDKGYSYNIR YKISPGGISQ IKKEPIQPET 2 ID NO:69 PDM3 to start and stop c 41 GGAAAAGGCT AAACCACATG CACCAGAGAT CGCTGGAAAT TGCCGTGATT CATACTGGAG AACCTCCTTA GGGAAAGGCT	HSPGKEGKRT ILDLVKGTHY PQPKPSVQKT DNA SEQUENCE bdons) 51 TCATCAAGAA GATGCAGCCT CTCATACAGG CACAGCTCAA GTGGAAAAGG AAAAACCCTA ATCATCGACG TCAGCCAGAA	360 420 480 60 120 180 240 300 360 420 480
45 50 55	DYTPFSCIKI CVACQKYFEM KDASSALASL Nucleic Acid Acce Coding sequence: 1	LFLKGIGLTL ILSNPPSGGD IHNVDDCGFS NSSLEHDMEG SSION #: NM_02 108-491 11	EQALQFWKQE YHGCPFRHSD LNHPNQFFCE LEDYFSEDS 4840 (underlined sequ 21 CATATATATA AGAGATTCA AAAGGTTCAA CAGGAGAGAA TCATTGTACA AAAGGTTCAC AAAGGTTCAC AAAGGTTCAC AAAGGTTCAC AAAGGTTCAC AAAGGTTCAC AAAGGTTCAC AAAGGTTCAC	FIKGKMDPDK FELLKQKLQS SQRILNGGKD SEC ences correspond 31	FDKGYSYNIR YKISPGGISQ IKKEPIQPET 2 ID NO:69 PDM3 to start and stop c 41 GGAAAAGGCT AAACCACATG CACCAGGAA TGCCTGGAAT TGCCGTGATT CATACTGGAG AACCTCCTTA GGGAAAAGGCT ACACCTTTG	HSPGKEGKRT ILDLVKGTHY PQPKPSVQKT DNA SEQUENCE codons) 51 TCATCAAGAA GATGCAGCCT CTCATACAGG CACAGCTCAA GTGGAAAAGG AAAACCCTA TTCATCACGAG TCAGCCAGAA TATCTACTGA	60 120 180 240 300 360 420 540
45	DYTPFSCIKI CVACQKYFEM KDASSALASL Nucleic Acid Acce Coding sequence: 1 AATTCATACA GTCTCGGCTC GTGTGGGAAG AGAGAAGCCCC TGCACATCAG TATATCAGAT TACTCACACT GACATGTTAG GACATGTTAG GTGTGGAAAA	LFLKGIGLTL ILSNPFSQGD IHNVDDCGFS NSSLEHDMEG SSION #: NM_02 108-491 11	EQALQFWKQE YHGCPFRHSD LNHPNQFFCE LEDYFSEDS A840 (underlined sequ CATATATATA AGAGATTCA AAAGGTCAG CTGAATGTGA TCATTGTACA AAGGCTTCAT CATATGATA AGAGATTCA AAGGTTCAT CATATGTACA AAGGTTCAT CATATGTACA AAGGTTCAT AAAGTTCAG	FIKGKMDPDK FELLKQKLQS SQRILNGGKD SEC ences correspond 31 CAGTGATTGT TACAGGAGAG GCTCACTGAA CAAAGCATTC GTCATATATA TCAGCGAATT CCAAAAGGGC CAAATGAATG CAATGAATG CAAAGGAT CACAGGAAAG	FDKGYSYNIR YKISPGGISQ IKKEPIQPET 2 ID NO:69 PDM3 to start and stop c 41 GGAAAAGGCT AAACCACATG CACCAGAGAA CGCTGGAAAT TGCCGTGATT CATACTGGAG AACCTCTTA GGGAAAGGCT ACACCCTTTG ACACCCTTTG CACACCAGAGAA	HSPGKEGKRT ILDLVKGTHY PQPKPSVQKT DNA SEQUENCE odons) 51 TCATCAAGAA GATGCAGCCT CTCATACAGG CACAGCTCAA GTGGAAAAGG AAAACCCTA TCATCGACG TCAGCCAGA TTCATCGACG TCAGCCAGAA TATGTACTGA TTCATCACAG TTCACACAGG TTCACACAGG TTCACACAGG	60 120 180 240 360 420 480 540 600
45 50 55	DYTPFSCIKI CVACQKYFEM KDASSALASL Nucleic Acid Acce Coding sequence: 1 AATTCATACA GTCTCGGCTC GTGTGGGAAG AGAGAAGCCC TGCACATCAG TATATCAGAT TACTCACACT GACATCAG GACATCTTTAG GTGTGGAAAA AGAGAAACCC	LFLKGIGLTL ILSNPFSGGD ILNNVDDCGFS NSSLEMDMEG SSION #: NM_07 108-491 11	EQALQFWKQE YHGCPFRHSD LINHPNOFFCE LEDYFSEDS 4840 (underlined sequ 21 CATATATATA AGAGATTCA AAAGGTCCA CTGAATGTGA TCATTGTACA AAGGCTTCAT AGAGATTCA AGAGATTCA AGAGATTTCA AGAGTTTCAG	FIKGKMDPDK PELLKQKLQS SQRILNGGKD SEC ences correspond 31	FDKGYSYNIR YKISPGGISQ IKKEPIQPET ID NO:69 PDM3 to start and stop c 41	HSPGKEGKRT ILDLVKGTHY PQPKPSVQKT DNA SEQUENCE odons) 51 TCATCAAGAA GATGCAGCCT CTCATACAGG CACAGCTCAA GTGGAAAAGC AAAACCCTA TTCATCGACG TCAGCCAGAA TATGTACTGA TATGTACTGA CATGTCCAGG CATGTCTCAA	360 420 480 600 120 180 240 300 420 480 540 600 660
45 50 55	DYTPFSCIKI CVACQKYFEM KDASSALASL Nucleic Acid Acce Coding sequence: 1 AATTCATACA GTCTCGGCTC GTGTGGGAAG AGAGAAGCCC TCACATCAG TATATCAAT TACTCACACT GACATCTAG GTGTGGAAAA AGAGAACCC CCAGACATCG	LFLKGIGLTL ILSNPPSGGS NSSLEHDMEG SSION #: NM_02 108-491 11	EQALQFWKQE YHGCPFRHSD LNHPNQFFCE LEDYFSEDS 4840 (underlined sequ 21 CATATATATA AGAGATTCA AAAGGTCCAA AAGGTCTCAA AGGCTTCAT CCTATATATATA AGAGATTCAA AGAGATTCAA AGAGATTCAA AGAGATTCAA AGAGATTCAA AGAGATTCAA ACAACTCAGG AGAGATTTCA ACAACTCAGG AGAGATTTCA ACAACTCAGG AGAGCTTCAGG CGGACTAGGG CGGACTAGGG CAGGGGGAGAG	FIKGKMDPDK FELLKQKLQS SQRILNGGKD SEC ENCES COTESPOND 31	FDKGYSYNIR YKISPGGISQ IKKEPIQPET ID NO:69 PDM3 to start and stop c 41	HSPGKEGKRT ILDLVKGTHY PQPKPSVQKT DNA SEQUENCE coons) 51 TCATCAAGAA GATGCAGCCT CTCATACAGG CACAGCTCAA GTGGAAAAGC TATCATCACGA TCATCACAC TCACCAGAA TATGTACTGA TTCACACAGG CATGTCTCAA GTGGAAAGC TTCACACAGG CATGTCTCAA GTGGGAAAGC	60 120 120 130 360 420 360 420 540 600 720
45 50 55	DYTPFSCIKI CVACQKYFEM KDASSALASL Nucleic Acid Acce Coding sequence: 1	LFLKGIGLTL ILSNPPSGGD IHNVDDCGFS NSSLEHDMEG SSION #: NM_02 108-491 11 GGAGAGAAGT ATTAATCATC GGCTTCTCCA AAAGCTCACA AAAACTCACA AAAACTCATA TCTCATCCAC TCTGCTCAC TATACATGCA AGAACTCATA TTGTCATGCC	EQALQFWKQE YHGCPFRHSD LNHPNQFFCE LEDYFSEDS 4840 (underlined sequ 21 CATATATATO AGAGATTCA AAAGGTTCA CAGGAGAGA TCATTGTACA AAGGCTTCAT AGAGATTCAC AGAGTTCAC GTGACTGGG CAGGGGAGAG TTGTTTATCA	FIKGKMDPDK FELLKQKLQS SQRILNGGKD SEC ences correspond 31	FDKGYSYNIR YKISPGGISQ IKKEPIQPET 2 ID NO:69 PDM3 to start and stop c 41 GGAAAAGGCT AAACCACATG CACCAGAAA CGCTGGAAAT CGTGGAAAT CATACTGGAG AACCTCTTAG GGGAAAGGCT ACACCTTTG CACCAGAGAA AGAACACATTG CACCAGAGAA TGCTCTGATT TGCTCTGATT CCTCCATTCATT CTGCATGCATC	HSPGKEGKRT ILDLVKGTHY PQPKPSVQKT DNA SEQUENCE codons) 51 TCATCAAGAA GATGCAGCCT CTCATACAGG CACAGCTCAA GTGGAAAAGG AAAACCCTA TTCATCGACG TCAGCCAGAA TATCATCGACG TCAGCAGAA TATCTACTGA TTCACACAGG CATGTTCTCAA GTGGAAAAG GTGGGAAAGC GAGAGAATG	60 120 180 240 300 360 420 480 540 660 720 780
45 50 55 60	DYTPFSCIKI CVACQKYFEM KDASSALASL Nucleic Acid Acce Coding sequence: 1	LFLKGIGLTL ILSNPPSGGD IHNVDDCGFS NSSLEHDMEG SSION #: NM_02 108-491 11 GGAGAGAAGT ATTAATCATC GGCTTCTCCA AAAGCTCACA AAAACTCACA AAAACTCATA TCTCATCCAC TCTGCTCAC TATACATGCA AGAACTCATA TTGTCATGCC	EQALQFWKQE YHGCPFRHSD LNHPNQFFCE LEDYFSEDS 4840 (underlined sequ 21 CATATATATO AGAGATTCA AAAGGTTCA CAGGAGAGA TCATTGTACA AAGGCTTCAT AGAGATTCAC AGAGTTCAC GTGACTGGG CAGGGGAGAG TTGTTTATCA	FIKGKMDPDK FELLKQKLQS SQRILNGGKD SEC ENCES COTESPOND 31	FDKGYSYNIR YKISPGGISQ IKKEPIQPET 2 ID NO:69 PDM3 to start and stop c 41 GGAAAAGGCT AAACCACATG CACCAGAAA CGCTGGAAAT CGTGGAAAT CATACTGGAG AACCTCTTAG GGGAAAGGCT ACACCTTTG CACCAGAGAA AGAACACATTG CACCAGAGAA TGCTCTGATT TGCTCTGATT CCTCCATTCATT CTGCATGCATC	HSPGKEGKRT ILDLVKGTHY PQPKPSVQKT DNA SEQUENCE codons) 51 TCATCAAGAA GATGCAGCCT CTCATACAGG CACAGCTCAA GTGGAAAAGG AAAACCCTA TTCATCGACG TCAGCCAGAA TATCATCGACG TCAGCAGAA TATCTACTGA TTCACACAGG CATGTTCTCAA GTGGAAAAG GTGGGAAAGC GAGAGAATG	60 120 120 130 360 420 360 420 540 600 720
45 50 55	DYTPFSCIKI CVACQKYFEM KDASSALASL Nucleic Acid Acce Coding sequence: 1 AATTCATACA GTCTCGGCTC GTGTGGGAAG AGAGAAGCCC TGCACATCAG TATATCAG TATATCAG TATATCAG TATATCAG TATATCAG TATATCAG TATATCAG TATATCAGAT TACTCACACT GACATGTTA AGAGAAACCC CAGACATCG TTTCTCCCAC TGTAGGTTCA	LFLKGIGLTL ILSNPFSQGD IHNVDDCGFS NSSLEHDMEG SSION #: NM_02 108-491 11 GGAGAGAAGT ATTAATCATC AAAGGTCACA AAAGGTCACA AAAGGTCACA AAAGTCACA ATATCACTCAC TCCTGCTCAC TCCTGCTCAC TATACATGCA AGAACTCATC TGTGATGCA AGACTCATC TGTGATGCA AGACTCATC TGTGATGCA AGACTCATC TGTGATGCA AGACTCATC TGTGATGCA AGACTCATC TGTGATGCA AGACTCATC TGTGCTCATC TGTGATTGGC	EQALQFWKQE YHGCPFRHSD LNHPNQFFCE LEDYFSEDS A840 (underlined sequ 21 CATATATATO AGAGATTCA AAAGGTCAGA CTGAATGTACA AAAGGTTCAT CATATATACA AAAGGTTCAT CATATATACA ACAGTCAGT CATATATACA ACAGTCAGT CATATGAATTCA ACAGTCAGT CATATGAATTCA ACAGTCAGT CATATGAATTCA ACAGTCAGT AGAGATTCAA ACAGTCAGG GTGACTTGAGT CAGGGGAGAGA AAAATCCTTG	FIKGKMDPDK FELLKQKLQS SQRILNGGKD SEC ences correspond 31	FDKGYSYNIR YKISPGGISQ IKKEPIQPET ID NO:69 PDM3 to start and stop c 41 GGARAAGGCT AAACCACATG CACCAGGAA CGCTGGAAAT TGCCGTGATT CATACTGGAG AACCTCTTA GGGAAAGGCT ACACCCTTTG CACCAGAGAA TGCTCTGAT TGCTCTGAT TGCTCTGAT TGCTCTGAT CTGCATGCAA CATAGCTTAT	HSPGKEGKRT ILDLVKGTHY PQPKPSVQKT DNA SEQUENCE odons) 51 TCATCAAGAA GATGCAGCCTA CTCATACAGG CACAGCTCAA GTGGAAAGG AAAACCCTA TTCATCGACG TCAGCCAGAA TATGTACTGAC TCACACAGG CATGTCTCAA GTGGGAAAGC CAGGTAAAGC CAGGAAATG CACATACACG CACATACACG	60 120 180 240 300 360 420 480 540 660 720 780
45 50 55 60	DYTPFSCIKI CVACQKYFEM KDASSALASL Nucleic Acid Acce Coding sequence: 1	LFLKGIGLTL ILSNPFSGGD IHNVDDCGFS NSSLEMDMEG SSION #: NM_02 108-491 11 GGAGAGAAGT ATTAATCATC GGCTTCTCCA TATGAATGCA AAGGGAAACT AAGGGAAACA GAAGTCACA GGAGGAAACA TATGCATCCA TCTGCTCCA TATACATGCA AGAACTCATA TTGTCATGCA GGTCAAATGG CGCATAAAAG CAGGATAAAG	EQALQFWKQE YHGCPFRHSD YHGCPFRHSD ARAO (underlined sequ 21 CATATATATA AGAGATCAG CAGAGAGAA TCATTGTAG AGAGATTCAA AGAGCTTCAT AGAGATTCAT AGAGATTTAA AGAGATTTAA AGAGATTTAA AGAGATTTAA ACAGTCAGG GTGACTGTTAACA ACAGTCAGG AGAGATTTAACA ACAGTCAGG AGAGATTTCAA ACAGTCAGG ACAGTCAG	FIKGKMDPDK FELLKOKLOS SQRILNGGKD SEC ences correspond 31	FDKGYSYNIR YKISPGGISQ IKKEPIQPET ID NO:69 PDM3 to start and stop c 41	HSFGKEGKRT ILDLVKGTHY PQPKPSVQKT DNA SEQUENCE odons) 51 TCATCAAGAA GATGCAGCCT CTCATACAGG AAAAACCCTA TTCATCGACG TCAGCCAGA TATGTACTGA TTCATCGACG CATGTCTCAA GTGGAAAGC CATGTCTCAA GTGGAAACC CATGTCTCAA CTGGGAAACC CATGTCTCAA CTGGGAAACC CACATACACG CTCTGTGGC CTTCTGTGCC	360 420 480 60 120 180 360 360 420 480 540 660 720 780 840 900
45 50 55 60	DYTPFSCIKI CVACQKYFEM KDASSALASL Nucleic Acid Acce Coding sequence: 1	LFLKGIGLTL ILSNPPSGGD IHNVDDCGFS NSSLEHDMEG SSION #: NM_00 108-491 11	EQALQFWKQE YHGCPFRHSD LNHPNQFFCE LEDYFSEDS 4840 (underlined sequ 21	FIKGKMDPDK FELLKQKLQS SQRILNGGKD SEC ENCES COTTESPOND 31	FDKGYSYNIR YKISPGGISQ IKKEPIQPET ID NO:69 PDM3 to start and stop c 41	HSPGKEGKRT ILDLVKGTHY PQPKPSVQKT DNA SEQUENCE coons) 51 TCATCAAGAA GATGCAGCCT CTCATACAGG CACAGCTCAA GTGGAAAAGC TATCATCACAGA TATGTACTGA TTCACACAGA CATGTCTCAA GTGGAAAACC CACATACACG CACATACACG CTTCTGTGGC CCATTGTGAC CCATTGTGAC	60 120 120 120 130 360 420 540 660 720 780 840 900 960
45 50 55 60	DYTPFSCIKI CVACQKYFEM KDASSALASL Nucleic Acid Acce Coding sequence: AATTCATACA GTCTCGGCTC GTGTGGGAAG AGAGAAGCC CTTCATTCAG TATATGCAAT TACTCACAT TACTCACAT GACATGTTA GTGTGGAAAA AGAGAAACCC CAGACATCG TTTCTCCCAC TGTAGGTTCA TGACTTCATA AGCTCAGAC CCAGCCTGTT	LFLKGIGLTL ILSNPPSGED IHNVDDCGFS NSSLEHDMEG SSION #: NM_OZ 108-491 11 GGAGAGAGAGT ATTAATCATC GCCTTCTCCA AAAGCTCACA AAAGCTCACA AAAGCTCACA AAAGCTCACA TATACATGCA TCTGCTCAC TATACATGCA TGGATAACTG TGTCATGCC GTCAAATTGG CAGGATAAACT TCGCAAATTGG CAGGATAACTG TCATACATACATGCC GTCAAATTGG CAGGATAACTG TCATTACATGCC GTCAAATTAGCT CAGGATAAACTA TCGCAGAAGTT GCCAGAAGTT	EQALQFWKQE YHGCPFRHSD LNHPNQFFCE LEDYFSEDS 4840 (underlined sequ 21 CATATATATO AGAGATTCA AAAGGTTCA AAAGGTTCA ACAGTCAGA GTGATTGAA ACAGTCAGG GTGACTTGGA CAGGGGAGAG TTGTTTATCA AAAATCCTTG ACTGTTTACA ACAGTCTTGA ACTCTGTTGA ACAGTCGGTT CAGTCTCAGG	FIKGKMDPDK FELLKQKLQS SQRILNGGKD SEC ences correspond 31	FDKGYSYNIR YKISPGGISQ IKKEPIQPET ID NO:69 PDM3 to start and stop c 41 GGARARAGGCT ARACCACATG CACCAGARAA CGCTGGARAT CATACTGGAG ARACTCTTA GGGARAGGCT ACACCATTG CACCAGAGAA TGCCTGATT CTGCAGAGAA CGATAGATAA CTTGCATTGAT CTGCATGATA CTGCATGATA CTGCATGATA CTGCATGATA CATAGCTTAT CTGCAGAGAA CATAGCTTAT CTGCAGAGAA CATAGCTTAT	HSPGKEGKRT ILDLVKGTHY PQPKPSVQKT DNA SEQUENCE codons) 51 TCATCAAGAA GATGCAGCTCAA GATGCAACACG CACAGCTCAA GTGGAAAAGG AAAACCCTA TTCATCGACG TCAGCAGAA TATGTACTGA GTGGAAAAG CATGTCTCAA GTGGAAAAC CAGAGAAATG CACATACACG CATTCTGTGGC CATTCTGTGGA AATAAAAACC	60 120 180 240 300 360 420 600 660 720 840 900 1020
45 50 55 60	DYTPFSCIKI CVACQKYFEM KDASSALASL Nucleic Acid Acce Coding sequence: 1 AATTCATACA GTCTCGGCTC GTGTGGGAAG AGAGAAGCCC TGCACATCAG TATATCAGATT GACATTTAG GTGTGGAAAA AGAGAAACCC CAGACATCGG TTTCTCCCAC TGTAGGTTCA TGATCTCATA AGCTCAGAC CCAGCCTGTT ATATGAATCC CCAGCCTGTT ATATGAATCC CAGCCTGTT ATATGAATCC	LFLKGIGLTL ILSNFPSGGD ILNIVDDCGFS NSSLEMDMEG SSION #: NM_07 108-491 11	EQALQFWKQE YHGCFFRHSD YHGCFFRHSD LINHPNOFFCE LEDYFSEDS 4840 (underlined sequ 21 CATATATATATA AGAGAGTTCA AGAGGAGGAA TCATTGTACA AAAGGCTTCAT CCTATGAATG AGAGGTTCAT CCTATGAATG AGAGGTTCAT CCTATGAATG AGAGGTTCAT AGAGGTTTGAT ACAGTGCGTT ACAGTGCGTT ACAGTGCGTTT ACAGTGCGTTT CAGTCTCAGG GTACTTCAGG GTACTTCTTTTAA	FIKGKMDPDK FELLKQKLQS SQRILNGGKD SEC ences correspond 31	FDKGYSYNIR YKISPGGISQ IKKEPIQPET ID NO:69 PDM3 to start and stop c 41	HSPGKEGKRT ILDLVKGTHY PQPKPSVQKT DNA SEQUENCE odons) 51 TCATCAAGAA GATGCAGCCT CTCATACAGGA ATAACCCTA TTCATCGACG TCAGCCAGAA TATGTACTGA TTCATCGACG CATGTCTCAA GTGGGAAAGC CATGTCTCAA GTGGGAAAGC CATGTCTCAA GTGGGAAAGC CACATACACG CACATACACG CACATACACG CTTCTGTGGC CACATACACAC CTTCTGTGGC CACATACACAC TGTCACAAAA	60 120 180 240 300 360 420 480 540 660 720 780 840 900 960 1020 1080
45 50 55 60	DYTPFSCIKI CVACQKYFEM KDASSALASL Nucleic Acid Acce Coding sequence: 1	LFLKGIGLTL ILSNPFSGGS ILSNPFSGGS NSSLEMDMEG SSION #: NIM_OZ 108-491 11 GGAGAGAAGT ATTAATCATCA GAACTCACA AAAGCTCACA AAAGCTCACA AAAGCTCACA AAACTCATCA GGAGAAAACT CGTCACATA AGAACTCATA TTGTCATGCA CGTCAAATTGG CAGGATAAAG TCATTAACTA GCCAGAAGTT AGTGAATTGG AACTGAATTGG AACAAACTGA	EQALQFWKQE YHGCPFRHSD YHGCPFRHSD LENHPNQFFCE LEDYFSEDS 4840 (underlined sequ 21 CATATATATA AGAGATTCA AAAGGTCCAB CTGAATGTGA CAGGAGAGAA TCATTGTACA AAGGTTCAT AGAGTTTAT AGAGTTTAT AGAGTTTAT ACAGTGGGG AAAATCTTG AAAATCCTTG AAAATCCTTG ACTCTGTTAA ACAGTGCGTT TATATTCAAG CTAGGTGCTTT TATATTCAAG	FIKGKMDPDK FELLKOKLOS SQRILNGGKD SEL ENCES COTESPOND 31	FDKGYSYNIR YKISPGGISQ IKKEPIQPET 2 ID NO:69 PDM3 to start and stop of the start and start a	HSPGKEGKRT ILDLVKGTHY PQPKPSVQKT DNA SEQUENCE bdons) 51 TCATCAAGAA GATGCAGCCTA CATACAGG CACAGCTCAA ATGCACCTAA TTCATCGACG TCAGCCAGAA TATGTACTGA TTCACACAGG CATGTCTCAA GTGGAAAAGC CACATACACG CACATACACG CTTCTGTGGC CATTGTGGC CATTGTGAG AATAAAAACC TTCTCAAAA ACCTTATGGC	360 420 480 60 120 180 360 360 420 480 540 660 720 780 900 960 1020 1080 1140
45 50 55 60	DYTPFSCIKI CVACQKYFEM KDASSALASL Nucleic Acid Acce Coding sequence: 1	LFLKGIGLTL ILSNPPSGGD LHNVDDCGFS NSSLEHDMEG SSION #: NM_00 108-491 11	EQALQFWKQE YHGCPFRHSD LNHPNQFFCE LEDYFSEDS 4840 (underlined sequ 21 CATATATATAT AGAGAGTTCA AAAGGTCCAA AAGGTCTAA AGAGTTCAA AGAGTTCAA AGAGTTCAA AGAGTTCAA AGAGTTCAA AGAATTCAAG ACAAGTCAGG CTGAATGTAAT ACAAGTCAGG CTGACTGTGG CAGGGGAGAG TTGTTTATCA AAAATCCTTG ACTGTTTAA ACAGTGCGTT CAGTGTCAGG CTGATTCAGG ACTCTCTTAA ACAGTGCGTT CAGTCTCAGG ATATTCAAG AAAATTAGTA	FIKGKMDPDK FELLKQKLQS SQRILNGGKD SEC ENCES COTTESPOND 31	FDKGYSYNIR YKISPGGISQ IKKEPIQPET ID NO:69 PDM3 to start and stop of the start and	HSPGKEGKRT ILDLVKGTHY PQPKPSVQKT DNA SEQUENCE coons) 51 TCATCAAGAA GATGCAGCCT CTCATACAGG CACAGCTCAA ATTCATCGACG TCAGCAGAA TATGTACTGA TTCACACAGG CATGTCTCAA CAGGAAAACC CACATACACG CTTCTTGGC CCATTGTGGC ATTAAAAACC TGTCACAAGC ATTAAAAACC TGTCACAGAG ATTAAAAACC TGTCACAGAG ATTAAAAACC TCTCACAGAG ATTAAAAACC TCTTCACAGG TCTTCACAAGG ATTAAAAACC TCTTCACAAGG TCTTTTATGG	60 120 180 360 420 180 360 420 780 840 960 1020 1020 1140 1200
45 50 55 60	DYTPFSCIKI CVACQKYFEM KDASSALASL Nucleic Acid Acce Coding sequence: 1	LFLKGIGLTL ILSNPPSGGD LHNVDDCGFS NSSLEHDMEG SSION #: NM_00 108-491 11	EQALQFWKQE YHGCPFRHSD LNHPNQFFCE LEDYFSEDS 4840 (underlined sequ 21 CATATATATAT AGAGAGTTCA AAAGGTCCAA AAGGTCTAA AGAGTTCAA AGAGTTCAA AGAGTTCAA AGAGTTCAA AGAGTTCAA AGAATTCAAG ACAAGTCAGG CTGAATGTAAT ACAAGTCAGG CTGACTGTGG CAGGGGAGAG TTGTTTATCA AAAATCCTTG ACTGTTTAA ACAGTGCGTT CAGTGTCAGG CTGATTCAGG ACTCTCTTAA ACAGTGCGTT CAGTCTCAGG ATATTCAAG AAAATTAGTA	FIKGKMDPDK FELLKQKLQS SQRILNGGKD SEC ENCES COTTESPOND 31	FDKGYSYNIR YKISPGGISQ IKKEPIQPET ID NO:69 PDM3 to start and stop of the start and	HSPGKEGKRT ILDLVKGTHY PQPKPSVQKT DNA SEQUENCE bdons) 51 TCATCAAGAA GATGCAGCCTA CATACAGG CACAGCTCAA ATGCACCTAA TTCATCGACG TCAGCCAGAA TATGTACTGA TTCACACAGG CATGTCTCAA GTGGAAAAGC CACATACACG CACATACACG CTTCTGTGGC CATTGTGGC CATTGTGAG AATAAAAACC TTCTCAAAA ACCTTATGGC	360 420 480 60 120 180 360 360 420 480 540 660 720 780 900 960 1020 1080 1140
45 50 55 60	DYTPFSCIKI CVACQKYFEM KDASSALASI Nucleic Acid Acce Coding sequence: 1	LFLKGIGLTL ILSNPPSGGD IHNVDDCGFS NSSLEHDMEG SSION #: NM_OZ 108-491 11 GGAGAGAAGT ATTAATCATC GCCTTCTCAA AAAGCTCACA AAAGCTCACA AAAGCTCACA AAACTCATC TCTGCTCAC TATACATGCA AGAACTCATA TTGTCATGCC GTCAAATTGG CAGGATAACT TCTGCTACC GTCAAATTGG CAGGATAACT AGCACAACT AATACTCGAG CTTCTCATCA	EQALQFWKQE YHGCPFRHSD LNHPNQFFCE LEDYFSEDS A840 (underlined sequ 21 CATATATATA AGAGATTCA AAAGGTTCAA AAAGGTTCAA AGAGTTCAA AGAGTTCAA AGAGTTCAA AGAGTTCAA AGAGTTCAA ACAGTCAGG GTGACTGTGA ACAGTCAGG CAGGGGAGAG TTGTTTATCA AAAATCCTTG ACTGTTTAA ACAGTGGGTT CAGTCTCAGG GTAGTGCTTT CAGTCTCAGG GTAGTGCTTT CAGTCTCAGG GTAGTGCTTT CAGTCTCAGG AAAAATAGTA ACAGTGCGTT CAGTCTCAGG GTAGTGCTTT TATATTCAAG AAAAATAGTA GTGACCATAGG	FIKGKMDPDK FELLKQKLQS SQRILNGGKD SEC ences correspond 31	FDKGYSYNIR YKISPGGISQ IKKEPIQPET ID NO:69 PDM3 to start and stop c 41 GGARAAGGCT AAACCACATG CACCAGAGAA CGCTGGAAAT CATACTGAG AACCTCTTA ACCACCTTTG CACCAGAGAA AGGATAAGT CTGCATGATT CTGCATGAT CTGCAGAGAA CATAGCTTAT CTGCAGATGA ACTTCTAATAA ATTTGCACAG ACTTGAATAGA ATTTGCACAG ACTTGAATAGA ACTTGCAATAG ACTTGAATAGA ACTTGCAATAGAA CTTGCAGATGAG ACTTGAATAGAA ACTTGCAATAGAA CTTGCAGATGAG ACTTGCAATAGAA ACTTGCAGAAGC TTACATTCAATAAAA ACTTGCAGAAGC CTTGAATAGAA CCTGCAGAAC CTGCAGAACT CTGCAGATGAG ACTTGCACATAG CTTGCATATAAAA ACTTGCAGAACT CTGCAGATGAG ACTGGAGAAT CAGTGAGCTT	HSPGKEGKRT ILDLVKGTHY PQPKPSVQKT DNA SEQUENCE codons) 51 TCATCAAGAA GATGCAGCTCAA GATGCAACCTA TTCATCACAGA TATCATCACAG TCACCAGAA TATCATCACAG CATGCACAGA CATGCACAGA CATGCTCACA GAGGAAATC CACATACACG CACTTCTCTGTGC CACTTGTGGA AATAAAAACC TTCTCGTAGA AATAAAAACC TTCTCACAAAA ACCTTATGGA ATTCACAAAA ACCTTATTGGA AATAAAAACC TTTATTGGA AATAGTTGGTA	60 120 180 360 420 180 360 420 780 840 960 1020 1020 1140 1200
45 50 55 60	DYTPFSCIKI CVACQKYFEM KDASSALASI Nucleic Acid Acce Coding sequence: 1 AATTCATACA GTCTCGGCTC GTGTGGGAAG AGAGAAGCCC TGCACATCAG TATATCAGATT GACATGTTA GACATGTTA AGAGAAACCC CAGACATCC TTTCTCCAC TGTAGGTTCA TGATCTCATA AGCTCAGAC TGATGCTCAT AGCTCAGAC TGATGCTCAG TGATGCTCAG TGATCTCAGA AGACACAC CCAGCCTGTT ATATGAATCC AACACACAG TAATAACAC GAAATATAAT GAAGATACAT	LFLKGIGLTL ILSNPFSGGD ILNNVDDCGFS NSSLENDMEG SSION #: NM_02 108-491 11 GGAGAGAAGT ATTAATCATC GAATGTGGAA AAGGGAAATC GAATGTGGAA TTGTCATCA TTGTCATGC GTCAAATTGG CAGGATAAAG CAGGAATTAACTA TGTCATGC CAGCAGAATT AGTCATACTA AGTCAAATTGG CAGGATTAACTA CAGCAGAAGTT AGTCAATGG AACAAACTGA ATACTCAGAA CCTTCTCATCA CAGTCATCGAA	EQALQFWKQE YHGCFFRHSD LNHPNOFFCE LEDYFSEDS 4840 (underlined sequ 21 CATATATATA AGAGATTCA AAAGTCCAG CTGAATGTGA CATGATGTACA AAAGTCCAG GTGACTTGTA ACAGTCAG ACAGTCAG ACAGTCAG CAGGGGAGAG AAAATCCTTG ACTCTTTATA ACAGTCGGT TGTTTATC AGTCTCTGG GTACTTCTGT ACTCTCTTA ACAGTCCAG CGTACTTTTATA ACAGTCCTTT TATATTCAAG AAAATAGTAA AAAATAGTAA AAAATAGTAA AAAATCCTTGT AAGTCCATAG	FIKGKMDPDK FELLKOKLOS SQRILNGGKD SEC ences correspond 31	FDKGYSYNIR YKISPGGISQ IKKEPIQPET IID NO:69 PDM3 to start and stop c 41	HSPGKEGKRT ILDLVKGTHY PQPKPSVQKT DNA SEQUENCE odons) 51 TCATCAAGAA CATGCAGCCTCAA GTGGAAAAGC TCAGCCAGA TATGTACTGA TCATCACAGG CAGGTCAA GTGGAAAGC TCAGCAGAA TATGTACTGA GTGGAAAGC CAGTCTCAA GTGGAAAGC CATGTCTCAA CTGGGAAAGC CACATACACG CATGTCTGAG CACATACACG CTTCTGTGGC CACATACACG TCTCACAAAA ACCTTATGGC TCTTTTTATGG CTTTTTTTTGGT GTATGGTAGGT ATAGTAGGGT ATAGTAGGT GGTATCAGGG	360 420 480 60 120 180 240 300 360 420 480 540 660 720 780 960 1020 1080 1140 1200 1320
45 50 55 60	DYTPFSCIKI CVACQKYFEM KDASSALASL Nucleic Acid Acce Coding sequence: 1	LFLKGIGLTL ILSNPFSGGS NSSLEMDMEG SSION #: NIM_OX 108-491 11 GGAGAGAAGT ATTARTCATC GCCTTCTCCA AAAGCTCACA AAAGCTCACA AAACTCATC GATTACATGCA AGACTCATA TTGTCATGCA GGAGAAGTT AGTGAATTGG CAGGATTAACTA GCCAGAAGTT AATGTCATACT ACTGAATTGG CAGGATTAACTA GCCAGAAGTT AATACTCAGAG CTTCTCATCA AATACTCAGAG CTTCTCATCAC AATACTGAA	EQALQFWKQE YHGCPFRHSD LNHPNQFFCE LEDYFSEDS 4840 (underlined sequ 21 CATATATATO AGAGACTICA AAAGGTCCAB CTGAATGTGA CAGGGGAGAA TCATTGTACA AAAGTCCAB CTGATTTATCA AGAGTTCAT ACAGTCAGG CTGATTTATCA AAAATCCTG AAAATCCTTG ACTCTGTTAA ACAGTCTGGT TTTTTAGAG AAAATCGTT TATATTCAAG AAAATAGTA AAAATCGTT TATATTCAAG AAAATAGTA AAAATAGTA AAAATAGTA AAAATAGTA AAAATAGTA AAAATAGTA AAAATAGTA AAAATAGTA AAAATAGTA AAAATCCTTGT TTTTTGGAAGA	FIKGKMDPDK FELLKOKLOS SQRILNGGKD SEL CRICES COTESPOND 31	FDKGYSYNIR YKISPGGISQ IKKEPIQPET 2 ID NO:69 PDM3 to start and stop of the start and start a	HSPGKEGKRT ILDLVKGTHY PQPKPSVQKT DNA SEQUENCE bdons) 51 TCATCAAGAA GATGCAGCCT CTCATACAGG CACAGCTCAA ATACTACGAG TCAGCCAGAA TATCTACTGAA GTGGAAAAG CATGTCTCAA GTGGAAAAC CTCTTTGTGGC CCATTGTGGC CCATTGTGGC TCTCACAAAA ACCTTATGGC TCTCACAAAA ACCTTATGGC TCTTCACAAAA ACCTTATGGC TCTTTATGG ATAGTTGGTA ATAGTTGGTA GTGATCAGGG ATACAGGG TCTTTTATGG ATAGTTGGTA GGTATCAGGG GGCAGGGTTG	360 420 480 60 120 180 360 420 540 660 720 780 840 900 960 1020 1020 1140 1200 1220 1320 1380
45 50 55 60 65 70	DYTPFSCIKI CVACQKYFEM KDASSALASL Nucleic Acid Acce Coding sequence: 1	LFLKGIGLTL ILSNPPSGGD IHNVDDCGFS NSSLEHDMEG SSION #: NM_00 108-491 11 GGAGAGAAGT ATTATCATC GCCTTCTCATC AAAGCTCACATC TCTGCTCACATC TCTGATATCATC GCAGATATGGA AGAACTCATA TTGTCATGCC GTCAAATTGG GTCAAATTGGC AACAACTGA ATTCTCATCA GGATAAACTGA ATTCTCATCA GGATAAACTGA ATTCTCATCA GGAGAAAC ATTCTGCCTT	EQALQFWKQE YHGCPFRHSD YHGCPFRHSD A840 (underlined sequ 21 CATATATATO AGAGASTICA AAAGGICCAA AAAGGICCAA AAGGITCAA AAAGTICAGG GTGACTGTGA ACATTGTACA AAAATCCTIGG ACTGTTAA ACAGTCCAGG ACTGTTAA ACAGTCCAGG TTGTTTATCAA ACAGTCCAGT CAGTCTCAGG GTAGTCCTTTT TATATTCAAG AAAATACTTT CAGTCTCAGG GTAGTCCTTTT TATATTCAAG AAAATACTT TATATTCAAG AAAATACTT GTGACCATAG AAAATACTT TTTTGAAGA AACTCCTTGTT TTTTTGAAGA AACTCCTAGAGA ATCCTCAGAGA	FIKGKMDPDK FELLKQKLQS SQRILNGGKD SEC CRCCC COMESPOND 31	FDKGYSYNIR YKISPGGISQ IKKEPIQPET ID NO:69 PDM3 to start and stop c 41	HSPGKEGKRT ILDLVKGTHY PQPKPSVQKT DNA SEQUENCE codens) 51 1 TCATCAAGAA GATGCAGCCT CTCATACAGG AAAAACCCTA TTCATCGACA TATGTACTGA TTCATCGACA GATGCAGAAATC CACATACAGG CATGTCTCGAA ACTTGTGGC CCTTTGTGGC CCATTGTGGA AATAAAACC TTCACAAAA ACCTTATGGAC AATAAAACC TCTCACAAAA ACCTTATGGAC ATTGTACAGG ATTGTACAGG CTTTTTTATGG ATTGTTATCGG ATTGTTATCGGC CTTTTTATGG ATTGTTATCAGG ATTGTTATCAGG CGCAGGGTTG CTTATGAAAAT	60 120 180 240 360 420 360 420 780 840 960 1020 1080 1020 1140 1200 1380 1440
45 50 55 60	DYTPFSCIKI CVACQKYFEM KDASSALASI Nucleic Acid Acce Coding sequence: 1	LFLKGIGLTL ILSNPFSGGD ILNNVDDCGFS NSSLEMDMEG SSION #: NM_OZ 108-491 11 GGAGAGAAGT GGATTATCATC GAATGTGGAA AAGGGAAATC GAATGTGGAA TTGTCATGC TCTGCTCACA TTGTATTGC TCTGCTCACA CGCATTACTACTA AGAACTCATA TTGTCATGC CGTCAAATTGG CAGAATTACTACTA AGCAAATTGG CAGAATTGG CAGAAGTT AGTGAATGTG AACAAACTGA ATACTCAGAA CTTCTTCATCA GGAGAGAAAC CTTTTCATCACAC ATTCTCCATC CTTCTTCATCACAC ATTCTTCATCACACACA	EQALQFWKQE YHGCFFRHSD YHGCFFRHSD LINHPNOFFCE LEDYFSEDS 4840 (underlined sequ 21 CATATATATATA AGAGAGTTCA CAGAGAGAAA TCATTGTACA AAAGGCTTCAT CCTATGAATG AGAGATTCAG GTGACTTGTACA ACAGTCCAG GTGACTTGTACA ACAGTCCAG GTGACTTGTTATA ACAGTCCAG ACAGTCCAG ACAGTCCAG GTGACTTCAG ACAGTCCAG ACAGTCCAG AAAATATGTA AAAATAGTA AAAATAGTA AAAATAGTA AAAATAGTA AAAATAGTA AAAATAGTA AAAATAGTA AAAATAGTA AAAATAGTA AAGTCCTTGT TTTTGGAAGA AACTCCTCAGAG AAATATGAA	FIKGKMDPDK FELLKOKLOS SQRILNGGKD SEC ences correspond 31	FDKGYSYNIR YKISPGGISQ IKKEPIQPET ID NO:69 PDM3 to start and stop c 41	HSPGKEGKRT ILDLVKGTHY PQPKPSVQKT DNA SEQUENCE odons) 51 TCATCAAGAA GATGCAGCCT CTCATACAGGA GTGGAAAAGC TCAGCCAGA TATGTACTGA TCATCAGGC GAGGAAAGC CATGTCTCAA GTGGAAAGC CATGTCTCAA GTGGAAAGC CATGTCTCAA GTGGAAAGC CATGTCTCAA GTGGAAAGC CACATGTCTGA GTGGAAAGC CACTTATGGC CACATGTGGC CACTTATAGC CTTTTTATGG TCTCACAAAA ACCTTATAGG GTATCAGGG GGCAGGGTTG GTATCAGGG GCCAGGGTTG CTATGAAAAT GCCCTGTGAA	60 120 180 240 300 360 420 720 780 840 900 1020 1080 1140 1260 1320 1320 1320 1440 1500
45 50 55 60 65 70	DYTPFSCIKI CVACQKYFEM KDASSALASL Nucleic Acid Acce Coding sequence: 1	LFLKGIGLTL ILSNPFSGGS ILSNPFSGGS NSSLEHDMEG SSION #: NIM_OZ 108-491 11 GGAGAGAAGT ATTAATCATC GGCTTCTCA TATGAATGCA AAGGGAAATC GAATGTGGAA AGACTCATA TTGTCATGC CTCTCAC CTCTCAC TATCAATTGG CAGAATTGG CAGAATTGG CAGAATTGG CAGAATTGG CAGAATTGG CAGAATTGG CAGAATTGG CAGAATTGG CACAAACTGA ATACTCAGCA ACAAACTGA ATACTCAGCA ATTCTCATC CATTTTCTCATC GATTAACTA GGAGAAACTGA ATTCTTCATC ACTTTCATC CATTTTCATC TTAGAGATTT	EQALQFWKQE YHGCPFRHSD YHGCPFRHSD A840 (underlined sequ 21 CATATATATG AGAGATTCA ACAGGACAGA AAAGGTCCAG AGAGATTTA AGAGATTCAA ACAGTCAG CTGAATGTAG CAGGAGAGA ACAGTCAG CAGGAGAGA ACAGTCAG CAGGAGAGA ACAGTCAG CAGGAGAGA ACAGTCAG CAGGAGAGA ACATTCTTA ACAGTCAGG CAGGAGAGA ATTTTATCAAG ACAGTCAGG CAGTCATTCTAC ACTCTTCTA ACAGTCCTTT TATATTCAAG AAAATAGTAA AAAATCTTG AAAATAGTAA AAAATAGTAA AAAATAGTAA AAAATAGTAA AAAATAGTAA AAGTCCTTGT TTTTTGGAAGA ATCCTCAGAAG AAAATTAGAA CGATCAGAAA	FIKGKMDPDK FELLKOKLOS SQRILNGGKD SEC SCRICKS CORESPOND 31 CAGTGATTGT CAGTGATTGT TACAGGAGAG GCTCACATGAA CAAAGCATTC CAAAAGGAATT CAAAAGGGA TCTCATTAAC GAAAGCTTTC ACCGTATGGA TAAGGAATG CACAGGAAGG TCTCATTAAC CAAGGAAGT CATGGTGACTAC CATGGTGACTAC CATGGTGACTAC CATGGTGACTAC CAGGAAGGC TGAAGTGGA GGAATACTAA CAGTGATCAA CAGTGATCAA CAGTGATCAA CAGTGATCAA CAGTGATCAA CAGTGATCAA CAGTGATCAA CAGTGAAACAC CCTTTGAAGG GGAATCATAA AGAACACAC TCTAACATCA	FDKGYSYNIR YKISPGGISQ IKKEPIQPET 2 ID NO:69 PDM3 to start and stop c 41	HSPGKEGKRT ILDLVKGTHY PQPKPSVQKT DNA SEQUENCE odons) 51 TCATCAAGAA GATGCAGCCT CTCATACAGG CACAGCTCAA GTGGAAAAGC TCAGCCAGA TATGTACTGA TCACCAGG CACAGCTCAA GTGGAAAACC CTCTGTGGC CACATACACG CACATACACG CTCTGTGGC CACATACACG CTCTTGTGGC CACATACACG CTTCTTGTGGC CACATACACG CTTCTTGTGGC CACATACACG CTTCTTGTGGC CACATACACG CTTCTTGTGGC CACATACACG CTTCTTGTGGC CACATACACG CTTCTTGTGGC CCATTGTGGC CTTTTTATGG ATAGTTCGTA ACCTTATGGAAAA CCTTATGGAAAA CCTTATGGAAAA CCTTATGGAAAAT CGCCTGTGAA GATAATATAC	360 420 480 60 120 120 1240 300 360 540 660 720 780 960 1020 1140 1260 1260 1320 1380 1440 1560
45 50 55 60 65 70	DYTPFSCIKI CVACQKYFEM KDASSALASL Nucleic Acid Acce Coding sequence: 1	LFLKGIGLTL ILSNPFSGGS ILSNPFSGGS NSSLEHDMEG SSION #: NIM_OZ 108-491 11 GGAGAGAAGT ATTAATCATC GGCTTCTCA TATGAATGCA AAGGGAAATC GAATGTGGAA AGACTCATA TTGTCATGC CTCTCAC CTCTCAC TATCAATTGG CAGAATTGG CAGAATTGG CAGAATTGG CAGAATTGG CAGAATTGG CAGAATTGG CAGAATTGG CAGAATTGG CACAAACTGA ATACTCAGCA ACAAACTGA ATACTCAGCA ATTCTCATC CATTTTCTCATC GATTAACTA GGAGAAACTGA ATTCTTCATC ACTTTCATC CATTTTCATC TTAGAGATTT	EQALQFWKQE YHGCPFRHSD YHGCPFRHSD A840 (underlined sequ 21 CATATATATG AGAGATTCA ACAGGACAGA AAAGGTCCAG AGAGATTTA AGAGATTCAA ACAGTCAG CTGAATGTAG CAGGAGAGA ACAGTCAG CAGGAGAGA ACAGTCAG CAGGAGAGA ACAGTCAG CAGGAGAGA ACAGTCAG CAGGAGAGA ACATTCTTA ACAGTCAGG CAGGAGAGA ATTTTATCAAG ACAGTCAGG CAGTCATTCTAC ACTCTTCTA ACAGTCCTTT TATATTCAAG AAAATAGTAA AAAATCTTG AAAATAGTAA AAAATAGTAA AAAATAGTAA AAAATAGTAA AAAATAGTAA AAGTCCTTGT TTTTTGGAAGA ATCCTCAGAAG AAAATTAGAA CGATCAGAAA	FIKGKMDPDK FELLKOKLOS SQRILNGGKD SEC ences correspond 31	FDKGYSYNIR YKISPGGISQ IKKEPIQPET 2 ID NO:69 PDM3 to start and stop c 41	HSPGKEGKRT ILDLVKGTHY PQPKPSVQKT DNA SEQUENCE odons) 51 TCATCAAGAA GATGCAGCCT CTCATACAGG CACAGCTCAA GTGGAAAAGC TCAGCCAGA TATGTACTGA TCACCAGG CACAGCTCAA GTGGAAAACC CTCTGTGGC CACATACACG CACATACACG CTCTGTGGC CACATACACG CTCTTGTGGC CACATACACG CTTCTTGTGGC CACATACACG CTTCTTGTGGC CACATACACG CTTCTTGTGGC CACATACACG CTTCTTGTGGC CACATACACG CTTCTTGTGGC CACATACACG CTTCTTGTGGC CCATTGTGGC CTTTTTATGG ATAGTTCGTA ACCTTATGGAAAA CCTTATGGAAAA CCTTATGGAAAA CCTTATGGAAAAT CGCCTGTGAA GATAATATAC	60 120 180 240 300 360 420 720 780 840 900 1020 1080 1140 1260 1320 1320 1320 1440 1500
45 50 55 60 65 70	DYTPFSCIKI CVACQKYFEM KDASSALASL Nucleic Acid Acce Coding sequence: 1	LPLKGIGLTL ILSNPPSGGD INNVDCGFS NSSLEHDMEG SSION #: NIM_OX 108-491 11 GGAGAGAAGT ATTAATCATC GCCTTCTCCA AAAGCTCACA AAAGCTCACA AAATCCATC TCCTGCTATC GCAGATTAACTA GCAGAATTG GCAGAATTG GCAGAATTG CAGCAGAGTT AGTGAATTG ATTACAATGA ATACCATC TCTCACACA GACACTCACACA AGACTCATA TTGTCATCC CAGCAGTTT AGTGAATTG ACTAAACTGA ATACTCAGAG TTATCATCAC ACTACATGAA ATACTCAGAG TTCTCATCA ATACTCAGAG CTTCTCATCA AGACTATCAGAG CTTCTCATCA GGAGAAACTGAA GGAGAAACTGAA GGAGAAACTGAA TTCTGCCTT CATTTAGGAC TTTAGGACATTT TTTTAGGACATTT TTTTAGGACATTT TTTTAGGACATTT	EQALQFWKQE YHGCPFRHSD LNHPNQFFCE LEDYFSEDS 4840 (underlined sequ 21 CATATATATA AGAGACTICA AAAGGTCCAA CAGGAGAGAA TCATTCTACA ACAGTTCAA ACAGTTCAA ACAGTTCAA ACAGTTCAA ACAGTTCAA ACAGTTCAT CAGGAGAGAA ATATACCTTG ACAGTTCAT ACAGTCTCAT ACAGTCCTTT TATATTCAAG AAAATACTTG AAAATACTTG AAAATACTTG AAAATATTCAAG AAAATATTCAAG AAAATATTAAA ATATTCGAAGA AAGTCCTTGT TTTTTGGAAGA AACATTAGAA AACATTAGAA AACATTAGAA AACATTAGAA AACATTAGAA AACATTAGAA AACATTAGAA AACATTAGAA AATATACCTTG	FIKGKMDPDK FELLKOKLOS SQRILNGGKD SEC SCRICKS CORESPOND 31 CAGTGATTGT CAGTGATTGT TACAGGAGAG GCTCACATGAA CAAAGCATTC CAAAAGGAATT CAAAAGGGA TCTCATTAAC GAAAGCTTTC ACCGTATGGA TAAGGAATG CACAGGAAGG TCTCATTAAC CAAGGAAGT CATGGTGACTAC CATGGTGACTAC CATGGTGACTAC CATGGTGACTAC CAGGAAGGC TGAAGTGGA GGAATACTAA CAGTGATCAA CAGTGATCAA CAGTGATCAA CAGTGATCAA CAGTGATCAA CAGTGATCAA CAGTGATCAA CAGTGAAACAC CCTTTGAAGG GGAATCATAA AGAACACAC TCTAACATCA	FDKGYSYNIR YKISPGGISQ IKKEPIQPET ID NO:69 PDM3 to start and stop of the start and s	HSPGKEGKRT ILDLVKGTHY PQPKPSVQKT DNA SEQUENCE coons) 51 TCATCAAGAA GATGCAGCCTA CATCACAGG CACAGCTCAA CTGGAAAAGC TACACAGG CATGCAGAA TATCATCAGG CACAGCAGAA TATCATCAGG CACAGCAGAA TATCATCAGG CATTCTGAG CACATACAGG CTCTTGTGGC CTCTTCAGAAA ACCTTATGGGC TCTTTATGG ATTAGTTGGTA ATTAGTTGGTA ATTAGTTGGTA ATTAGTTGGTA CGCAGGGTTTG CTATGAAAA CGCTATCAGGG GCCAGGGTTTG CTATGAAAAT CCCCTGTGGA CATGAAAAT CCCCTGTGGA CATGATAATATAC ATGACTAATT	360 420 480 60 120 120 1240 300 360 540 660 720 780 960 1020 1140 1260 1260 1320 1380 1440 1560
45 50 55 60 65 70	DYTPFSCIKI CVACQKYFEM KDASSALASL Nucleic Acid Acce Coding sequence: 1	LFLKGIGLTL ILSNPPSGGD ILSNPPSGGD INNVDCGFS NSSLEHDMEG SSION #: NM_02 108-491 11	EQALQFWKQE YHGCPFRHSD YHGCPFRHSD LNHPNQFFCE LEDYFSEDS 4840 (underlined sequ 21 CATATATATO AGAGASTICA AAAGGICCAA AAAGGICCAA AAGGICTCAA AGAGTTCAA AGAGTTCAA AGAGTTCAA AGAATTCAGG GTGACTGTGA ACATGTCAGG GTGACTGTTAA ACAGTCCTGT TATATATCAAG ACAGTCCGTT TATATATCAAG AAAATCCTTGG GTACTCTTTAT TATATCAAG AAAATCCTTGT TATATTCAAG AAAATCCTTGT TATATTCAAG AAAATAGTA GTGACCATAG AAAATAGTA GTGACCATAG AAAATAGTA AGGTCCTTGT TTTTGGAAGA ATCCTCAGAG AAAATATGAA CGATCAGAGA AATCCTTGGAGAG AAAATATGAA CGATCAGAGAA ATCCTCAGAGG AAAATATGAA CGATCAGAGA AATCCTTGGAGAG AAAATATGAA CGATCAGAGAA ATCCTCAGAGG AAAATATGAA CGATCAGAGAA ATCCTTCGTCAGAGA AATCCTTGGAGAGAA ATCCTTCAGAGAA ATCATCAGAGAA ATCCTTCAGAGAGAA ATCCTTCAGAGAGAA ATCCTTCAGAGAGAA ATCCTTCAGAGAA ATCCTTCAGAGAGAA ATCCTTCAGAGAA ATCCTTCAGAGAAA ATCCTTCAGAGAAAA ATCCTTCAGAGAAAA ATCCTTCAGAGAAAA ATCCTTCAGAGAAAAAAAAAA	FIKGKMDPDK FELLKQKLQS SQRILNGGKD SEL ENCES COMESPOND 31	FDKGYSYNIR YKISPGGISQ IKKEPIQPET ID NO:69 PDM3 to start and stop c 41 GGARARAGGCT ARCCACATG CACCAGAGA TGCCTGGARAT GGGARAGGCT ARCCTCCTTG ARCCTCCTTG CACCAGAGAA AGGARARAT TGCTCTGATT CTGCAGGAT CTGCAGGAT AGCARATAA ACTGGAGAT ACTCTTGATTA CTGCAGGAT CTTGAATAA CTTGAATAA CTTGAATAA CTTGAATAA CTTGAATAA TTATATGGC TTACACCAGTA ACTGGAAATAA TTATATATGG TGAAATAAA TTATATATGC TGACCACATG TGAAATAAA TTATATATGC TGAAATAAA TTATATATGC TGAAATAAA TGAAATAAAA TTATATATGC TTACACCAATG	HSPGKEGKRT ILDLVKGTHY PQPKPSVQKT DNA SEQUENCE codons) 51 1 TCATCAAGAA GATGCAGCCT CTCATACAGG CACAGCTCAA TTCATCAGAG TTCATCAGAG TTCATCAGAG TTCATCAGAG TTCATCAGAG TTCACACAGG CATGTCTCAA ACCTTATGC CATTCTGAG CATTCTGAG ATTAAAAACC TTTCACAAAA ACCTTATGGC CCATTGTGGC CCATTGTGGAG AATAAAAACC TCTTATCAGG CTTCTTATCGG CTTCTTATCGG CTTTTTATGG ATTACAGGG GGCAGGGTTG CTATGAAAAT GCCTGTGAA GCTTATCAGGA GCCAGGGTTG CTATGAAAAT TACCTCTTCC	60 120 180 240 360 420 360 420 780 840 900 1020 1080 1260 1320 1260 1320 1440 1500 1500 1620 1680
45 50 55 60 65 70	DYTPFSCIKI CVACQKYFEM KDASSALASI Nucleic Acid Acce Coding sequence: 1	LFLKGIGLTL ILSNPFSGGD ILNNVDDCGFS NSSLEMDMEG SSION #: NM_02 108-491 11 GGAGAGAAGT GGATTATCATC GAATGTGGAA AAGGGAAAC ATATCCATC GTCAAATTGG CAGGATAAAG CAGGAATTAACTAA GCAGAAGT AACTCAGAA TTGTCATCC CATCATCACA ACAAACTGA ATACTCAGAA CTCATTCATCA CATCATCACA ACAAACTGA ATACTCAGAA CTCATTCATCA CATCATCACA C	EQALQFWKQE YHGCFFRHSD YHGCFFRHSD LINHPNOFFCE LEDYFSEDS 4840 (underlined sequ 21 CATATATATA AGAGAGTICA AAAGGTCCA CTGAATGTGA CAGGAGAGAA TCATTGTACA AAAGGTCCAG GTGACTGTGG CAGGGGAGAG AAAATCCTTG ACTCTGTTAA ACAGTCCAG GTACTGTTAA ACAGTCCTTT TATATTCAAG AAAATACTTT TATATTCAAG AAAATAGTA ATACTTGTTTTTTGGAAGA ATCCTTGT TTTTGGAAGA ATCCTCAGAG AAATATGTA CGATCAGAG AAATATGAA CGATCAGAG AAATATGAA CGATCAGAG AACTCCTTGT TTTTGGAAGA ATCCTCAGAG CACATCATGT TTTTGGAAGA ATCCTCAGAG CACATCATTGT TCTATTCCCA	FIKGKMDPDK FELLKOKLOS SQRILNGGKD SEC SCRICKS CORRESPOND 31 CAGTGATTGT CAGTGATTGT CAGTGATTGT CAAAGCATTC GTCATATATA CAAAGCATTC CAAAAGGATT CAAAAGGAT CAAAGGAATG CAAAGGAATG CAAAGGAATG CAAAGGAAAG TCTCATTAAC GAAAGCTTTC ACCGTATGGA TAAGGAATG CTCAGAGAGT CATGGAAAGG TCTAGAGAGG CTCTAGAGAG CTCTAGAGAG CTCTAGAGAG CTGAAATGT TCAGAAAACG CTTTGAAGG GGAATCATTA AGAACACAC TCTAACATCA AATCACTAG TCTAACATCA AATCACTAG TGTTAAAATTAA AGAACACAC TCTAACATCAA AATCACTAGT TGTTAAAATTAA AGAACACAC TCTAACATCAA AATCACTAGT TGTTAAAATTAA AGAACACAC TCTAACATCAA AATCACTAGT TGTTAAAATTAA AGAACACACAC TCTAACATCAA AATCACTAGT TGTTAAAATTAA ACCAAGATCAT ACCAAGATCAT CACAAGATCAT CACAAGATCAA ACCAAGATCAA ACCAAGATCA	FDKGYSYNIR YKISPGGISQ IKKEPIQPET IID NO:69 PDM3 to start and stop c 41	HSPGKEGKRT ILDLVKGTHY PQPKPSVQKT DNA SEQUENCE odons) 51 TCATCAAGAA CATGCAGCCT CTCATACAGGA GTGGAAAAGC TCAGCCTAA TTCATCGACG TCAGCCAGAA TATGTACTGA GTGGAAAGC CATGTCTCAA GTGGAAAGC CATGTCTCAA GTGGAAAGC CATGTCTCAA GTGGAAAGC CATGTCTCAA GTGGAAAGC CATGTCTGTGGC CATTGTGGC CATTGTGGC CATTGTGGC CATTGTGGG CATTAAAAAC CTTTTTATGG TTCTCTTTTATGG GTATCAGGG GGCAGGGTTG CTATGAAAA GCTTTTTATGG CTATGAAAA ACCTTATGGA AGTATCAGGG GGCAGGGTTG CTATGAAAAT TACCTCTTCC AGCTCTTTCC AGCTCTTTCTC	60 120 180 240 300 360 420 720 780 840 900 1020 1080 1140 1260 1320 1320 1440 1500 1680 1740
45 50 55 60 65 70	DYTPFSCIKI CVACQKYFEM KDASSALASI Nucleic Acid Acce Coding sequence: 1	LFLKGIGLTL ILSNPFSGGD ILNNVDDCGFS NSSLEMDMEG SSION #: NM_02 108-491 11 GGAGAGAAGT GGATTATCATC GAATGTGGAA AAGGGAAAC ATATCCATC GTCAAATTGG CAGGATAAAG CAGGAATTAACTAA GCAGAAGT AACTCAGAA TTGTCATCC CATCATCACA ACAAACTGA ATACTCAGAA CTCATTCATCA CATCATCACA ACAAACTGA ATACTCAGAA CTCATTCATCA CATCATCACA C	EQALQFWKQE YHGCFFRHSD YHGCFFRHSD LINHPNOFFCE LEDYFSEDS 4840 (underlined sequ 21 CATATATATA AGAGAGTICA AAAGGTCCA CTGAATGTGA CAGGAGAGAA TCATTGTACA AAAGGTCCAG GTGACTGTGG CAGGGGAGAG AAAATCCTTG ACTCTGTTAA ACAGTCCAG GTACTGTTAA ACAGTCCTTT TATATTCAAG AAAATACTTT TATATTCAAG AAAATAGTA ATACTTGTTTTTTGGAAGA ATCCTTGT TTTTGGAAGA ATCCTCAGAG AAATATGTA CGATCAGAG AAATATGAA CGATCAGAG AAATATGAA CGATCAGAG AACTCCTTGT TTTTGGAAGA ATCCTCAGAG CACATCATGT TTTTGGAAGA ATCCTCAGAG CACATCATTGT TCTATTCCCA	FIKGKMDPDK FELLKQKLQS SQRILNGGKD SEL ENCES COMESPOND 31	FDKGYSYNIR YKISPGGISQ IKKEPIQPET IID NO:69 PDM3 to start and stop c 41	HSPGKEGKRT ILDLVKGTHY PQPKPSVQKT DNA SEQUENCE odons) 51 TCATCAAGAA CATGCAGCCT CTCATACAGGA GTGGAAAAGC TCAGCCTAA TTCATCGACG TCAGCCAGAA TATGTACTGA GTGGAAAGC CATGTCTCAA GTGGAAAGC CATGTCTCAA GTGGAAAGC CATGTCTCAA GTGGAAAGC CATGTCTCAA GTGGAAAGC CATGTCTGTGGC CATTGTGGC CATTGTGGC CATTGTGGC CATTGTGGG CATTAAAAAC CTTTTTATGG TTCTCTTTTATGG GTATCAGGG GGCAGGGTTG CTATGAAAA GCTTTTTATGG CTATGAAAA ACCTTATGGA AGTATCAGGG GGCAGGGTTG CTATGAAAAT TACCTCTTCC AGCTCTTTCC AGCTCTTTCTC	60 120 180 240 360 420 360 420 780 840 900 1020 1080 1260 1320 1260 1320 1440 1500 1500 1620 1680

AAAATGTATT TAATTTAATA ATGTAACACA ACAAGTTTGG ATGTGTTTAA CTTTATAAAT 1860 AATCACCCCA GAGGAATGAA GTTCAAAACT TGTGAATAAC C

5	SEQ ID NO:70 P Protein Accession	DM3 Protein sequ n #: NP_0					
10			21 ELIQERSPMN EKNPIYAMNV			51 TQERSHIYAV NPMNAMNVGK	60 120
15		ession#: NM_0				DNA SEQUENCE	
	Coding sequence	: 341- 95	5 (underlined sequ	lences correspond	to start and stop (codons)	
20	1	11	21	31	41	51 	CO
20	AAGCCGACTG AGGTTGCACA CCCCCGTGCA	ACATAAGCCA CTTCTAAGAA GTCCCCTGTG	GCACAGTGAG GGTCCTAACG GAGCGGCGTG CCCAAGACAC	GAGCCTATGT GGGGGCTCGG AGCCTGATGC	GTAAGTCCAC CGACCTTCGC TTGTGCTCCG	TACTGGTGCA TTCAGTCGCT GTGGGCGGAC	120 180 240
25	GAGGAGTGAG GTTCATCAAG CTGGGATTTT	ACTGCAGGAG AGGACCATCT TTGTCTGAAA	AATTGGTGGC ATGTGGGCCG TGAAAATCCC ATCAACTGCA ATCTGTGTGA	TGCCAAAGAG CATGAATGAA GACTGTAAAT	ATGGATGAGA CTGACAACAA TTCCGACAGA	CTGTTGCTGA TCCTGAAGGC GAAAGGAATC	300 360 420 480 540
30	CCTGTTAGAC GATGAGTAAA GTTCAAGAAA GGAGAATGCA	ATCATTTATA GGACCAGGTG ATTCTTCAGA GTCTGGATTC	TGCAATTTCA AAGATGTTGA GAGCATTAAA GAATTGCCTG	TCAGCACCAG CCTTTTTGAT AAATGTGACA GGGAACACAG	AAAGTTTGGG ATGAAACAAT GTCAGCTTCA TACACAAAGC	ATGTTTTTCA TTAAAAATTC GAGAAACTGA CAAACCAGTA	600 660 720 780
35	GCTGAGGCGC CCGACAAGAG	AATACACCGC GAGATCATTT	ACTACTCCCA TTCTGGGTCA TAGATATTAC CCTCACAGTC	GGAGTTAGAA CGAAATGAAG	GCTACTGGGA AAAGCTTGCA	AAATCTACCT	840 900 960
40	SEQ ID NO:72 Protein Accession	DM8 Protein sequ n #: NP_00					
10	1	11 	21 	31 	41 	51 .	
45	ASISDAALLD VSFRETEENA	IIYMQFHQHQ	LTTILKAWDF KVWDVFQMSK YTKPNQYKPT KACN	GPGEDVDLFD	MKQFKNSFKK	ILQRALKNVT	60 120 180
	Shielala Aald Aaa	neeles # ND C		SE	Q ID NO:73 PDM9	DNA SEQUENCE	
50	Coding sequence	ession #: NM_0 : 1-1125	(underlined seque	ences correspond (to start and stop co	odons)	
	1	11	21	31	41	51	
	ATGGTGCTGT	GGGAGTCCCC	GCGGCAGTGC	AGCAGCTGGA	CACTTTGCGA	GGGCTTTTGC	60
55			CATGCTACTC				120 180
	GACAGAGAAA	ATGATCTCTT	CCTCTGTGAC	ACCAACACCT	GTAAATTTGA	TGGGGAATGT	240
			GACTTGCGTC GGAGAGCTAC				300 360
60	TGCAAACAGC	AGAGTGAGAT	ACTTGTGGTG	TCAGAAGGAT	CATGTGCCAC	AGATGCAGGA	420
			CCATGAAGGC TGGTGCAGAA				480 540
	GTGTGTAATA	TTGACTGTTC	TCAAACCAAC	TTCAATCCCC	TCTGCGCTTC	TGATGGGAAA	600
65			AATCAAAGAA TCAAGATAAC				660 720
	CATTATGCAA	GAACAGATTA	TGCAGAGAAT	GCTAACAAAT	TAGAAGAAAG	TGCCAGAGAA	780
			ACATTACAAT ATCTTGCAGG				840 900
70	GAAAAAAAGG	ACTACAGTGT	TCTATACGTT	GTTCCCGGTC	CTGTACGATT	TCAGTATGTC	960
10	ATCACAAGGA	AATGCCCCAG	AACAATTCAG AAGCAACAGA AAGAGCGTCC	ATTCACAGAC	AGAAGCAAAA		1020 1080

	SEQ ID NO:74 PDM9 Prot Protein Accession #:	ein sequence; NP_057276					
5	61 DRENDLFLCD	11 21 SSWTLCEGFC WL TNTCKFDGEC LR	LLLPVMLL IVAR IGDIVTCV CQFK	CNNDYV PVCGS	ENGESY QNECY	LRQAA	60 120
10	121 CKQQSEILVV 181 VCNIDCSQTN 241 HYARTDYAEN 301 EKKDYSVLYV 361 YSSDNTTRAS	FNPLCASDGK SY ANKLEESARE HH VPGPVRFQYV LI	DNACQIKE ASCQI IPCPEHYN GFCM	KQEKIE VMSLO HGKCEH SINMO	GROODN TTTTT	KSEDG TGQHC	180 240 300 360
15	Nucleic Acid Accession #: Coding sequence:		SE sequences correspond		DNA SEQUENCE		
20	1 11 	21 	31 	41	51 		
20	GGCGCCGGGA TTGGG TTCCTTCAGC GGGGC GTCCGGCCTG GCCCC GGTACGCGTG GACCG	ACTGG GAAGCGCC GGGCC GTNTCTGT	AT GCACTGCAG GC TATGGTCCTG	GGCATCTCGG GCTGACTTCG	TCGTGGAGCT GGGCGCGTGT	60 120 180 240	
25	CTCGCTAGTG CTGGA GGTCGGATGT GCTGC CAGAGATTCT GCAGC AGTTCAGGAA AGCTT TGTTCTCTCA AAAAT	TGGAG CCCTTCCG GGGAA AATCCAAG CTGCC GGTTAGCT TGGCA GAAGTGGT	CC GCGGTGTCAT GC TTATTTATGC GG CCACGATATC GA GAATCCGTAT	GGAGAAACTC CAGGCTGAGT AACTATTTGG GCCCCGCTGA	CAGCTGGGCC GGATTTGGCC CTTTGTCAGG ATCTCGTGGC	300 360 420 480 540	
30	TGACTITGCT GGTGG CACACGCACT GACAA AAGTICTITT CTGTG CATGTTGGAT GGTGG GGCTGTTGGA GCAAT	GGGTC AGGTCATT GAAAA CTCAGAAA AGCAC CTTTCTAT AGAAC CCCAGTTC	GA TGCAAATATG TC GAGTCTGTGG AC GACTTACAGG TA CGAGCTGCTG	GTGGAAGGAA GAAGCACCTC ACAGCAGATG ATCAAAGGAC	CAGCATATTT GAGGACAGAA GGGAATTCAT TTGGACTAAA	600 660 720 780 840 900	
35	GTCTGATGAA CTTCC TGCAGATGTA TTTGC TGCCTGTGTG ACTCC ACGGGGCTCG TTTAT GTTAAACACC CCAGC	AAAGA AGACGAAG GGTTC TGACTTTT CACCA GTGAGGAG CATCC CTTCTTCC	GC AGAGTGGTGT GA GGAGGTTGTT CA GGACGTGAGC AA AGGGGATCCT	CAAATCTTTG CATCATGATC CCCCGCCTTG TTCATAGGAG	ACGGCACAGA ACAACAAGGA CACCTCTGCT AACACACTGA	960 1020 1080 1140 1200	
40	GGAGATACTT GAAGA AATCATTGAA AGTAA AATTTGAATA CTGCA GAGGAACAGT ATTAC CTACAGTGAT GATTG	TAAGG TAAAAGCT TTTAC AGTGTAGA AGTGT CCTACCAC AATTC TAAAAATG	AG TCTCTAACTT GT AACACATAAC TC TAATCAAGAA GT TATCATTAGG	CCAGGCCCAC ATTGTATGCA AAGAATTACA GCTTTTGATT	GGCTCAAGTG TGGAAACATG GACTCTGATT TATAAAACTT	1260 1320 1380 1440	
45	TGGGTACTTA TACTA TGATATTAAG ATTCT TCTTGAAGAC ATCGA AAATGCCACA AATTG GGCCTTTTGT CTTGG	TGACT TATATTTT TATAC ATTTATTT TATGG TGATAAAA TGTTC ATGATCTC	GA ATGGGTTCTA AC ACTCTTGATT GT CACGTGAAAC CC TCTAAGCACA	GTGAAAAAGG CTACAATGTA AGAGTGATTG TTCCAAACTT	AATGATATAT GAAAATGAGG GTTGCATCCA TAGCAACAGT	1500 1560 1620 1680 1740	
50	TATCACACTT TGTAA CTGAAAAAA CATAT GGGACAGTCA GTTTT CTCTGGGCTG TCAGC TTCTGGATCT TATAC	CCAAA ATAATGAG AGGGT TGCCTGTA TTTCC TTTCTCCA CCAAC ACACAGCA	GA AATGTGTTGG TC CAGTAACTCG TG TGTTTGATTT AC ATCCAGAAAT	CTCACTACGT GGGCCTGTTT CTCCTCAGGC	AGAGTCCAGA CCCCGTGGGT TGGTAGCAAG	1800 1860 1920 1980 2040	
55	SEQ ID NO:76 PDO1 Protein Accession #:						
60	 1 MALQGISVVE	1 21 LSGLAPGRXC AM AVGCAAGALP PR GVLSKIGRSG EN	CHGETPAG PRDS	AAGKSK AYLC GGGLMC ALGI	QAEWIW PVQES	FCRLA KGQVI	60 120 180 240
65	241 YELLIKGLGL 301 EEVVHHDHNK 361 REEIYQLNSD	KSDELPNOMS TO ERGSFITSEE OD	DWPEMKKK FADVI VSPRLAPL LLNT:	PAKKTK AEWC(QIFDGT DACVT	PVLTP	300 360
70	Nucleic Acid Accession #: Coding sequence:		SE sequences correspond		DNA SEQUENCE codons)		
75	1 11 GTTAAATCCT TACTT					60	
80	CTTCACAGAG ACTTG AGAGTCAAAA TAGCT GCAGATTTGG ATCCA GCAAGGCATT ATACA TTGACTTCGG AACCT	GACAT GGGTTTTG GTAGT TGTGACAT AAGGC CATTGATA	CC AGATTATTCA TT TGGTATCGGG TA TGGGCAATAG	ATTCTCCTCT CTCCAGAACT GTTGTATATT	AAAGCCACTA TTTGCTTGGT TGCTGAATTG	120 180 240 300 360	

	CATCATGATC	AACTGGATCG	GATATTTAGT	GTCATGGGGT	TTCCTGCAGA	TAAAGACTGG	420
			AGAATATCCC				480
	TATGCCAACA	GTAGCCTCAT	AAAGTACATG	GAGAAACACA	AGGTCAAGCC	TGACAGCAAA	540
_	GTGTTCCTCT	TGCTTCAGAA	ACTCCTGACC	ATGGATCCAA	CCAAGAGAAT	TACCTCGGAG	600
5	CAAGCTCTGC	AGGATCCCTA	TTTTCAGGAG	GACCCTTTGC	CAACATTAGA	TGTATTTGCC	660
			CAAACGAGAA				720
			GCAGCAGAAC				780
			GGCGCCCCCA				840
10			CGGGGTCGGG				900
10			GCCTCCAAAC				960
			GCCCTCGGAT				1020
			CTCTCAGTCC				1080
			CCCATCTCAC				1140
15			AGAGCACAGG				1200
13			TGCCATTTAA TGATAGCTCT				1260 1320
			CCTGCATGTG				1320
			GTAACCTTCA				1440
			ACTATGGTTT				1500
20			CCAGTGGTGT				1560
20			TTTTAAAGCA				1620
			AACTGTTACT				1680
			GTCTAATAAA				1740
			TTTGCAGTTA				1800
25			ATTCATATGG				1860
			ATTACAATTT				1920
			ACAGTCTCGA				1980
			TAAGGAGTAA				2040
			GTATTGTACA				2100
30	TTCTTCACAT	CTGGGTCTGC	GTGAGTAACT	TTCTTGCATA	ATCAAGGTTA	CTCAAGTAGA	2160
	AGCCTGAAAA	TTAATCTGCT	AATAAAATTT	AGAGCAGTGT	TCTCCATTCG	TATTTGTATT	2220
	AGATATAGAG	TGACTATTTT	TAAAGCATGT	ATTTAAAAAT	GGTTTTATIC	ATGTTTAAAG	2280
			TTTTGCTGTT				2340
25			TTCTTTTGCC				2400
35			AAGACTATAC				2460
			TTCTTTCTTA				2520
			CTTAGAGCTC				2580
			CAGCACCAGC				2640
40			TGCAAATTAC				2700
40			CTGTGGGTCC				2760
			TTGATGTGAT				2820
			TAATGAAGTC				2880
			TTGGAAGGAG				2940 3000
45			ATTAGCAGCC CTCACTGTCC				3060
73			CCCTGCTGGG				3120
			TCTGTGACAG				3180
			GTCTTATAAT				3240
			CAAATCTGTT				3300
50			GCCAGTGATT				3360
			TTAACAACAA				3420
			TTGCAAGGAA				3480
			ACAAACCCAG				3540
			TTGTTAAATA				3600
55	CCTGTTTTCC	TTGTATATAA	TGACTTTTGC	TGGCAGAACT	GAAATATAAA	CTGTAAGGGG	3660
	ATTTCGTCAG	TIGCTCCCAG	TATACAATAT	CCTCCAGGAC	ATAGCCAGAA	ATCTCCATTC	3720
	CACACATGAC	TGAGTTCCTA	TCCCTGCACT	GGTACTGGCT	CTTTTCTCCT	CTTTCCTTGC	3780
			CTGATTCCCT				3840
~	CTTTCCTTTA	AAGGGGAACA	AAGCCTTTTT	TTTTTTTGAG	ACGGAGTGTT	GCTCTGTCAC	3900
60			ACGATCTTGG				3960
						CACCACGTCT	
						GCTGGTCTTG	4080
						ATTATAGGTG	
65						CCCTCAAACC	
65						TIGIGGICCC	
						AAGGGTACAG	
						ACAGGAAGGA	
						AGTTAGCTTT	
70						AGCTCAAGGT	
70						TGATGAAGTA	
						CIGIGGITIT	
						ACGAACATAA	
						ATTCCTGGTG	
75						GCTAAAGGTA	
13						TTTGAGTAAA AATCCAAAAC	
						TAGACCAACA	
						TTTAGGTCCC	
						TTTTTAAAAA	
80						GGGTGATATG	
							5205

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60	1	11	21	31	41	51				
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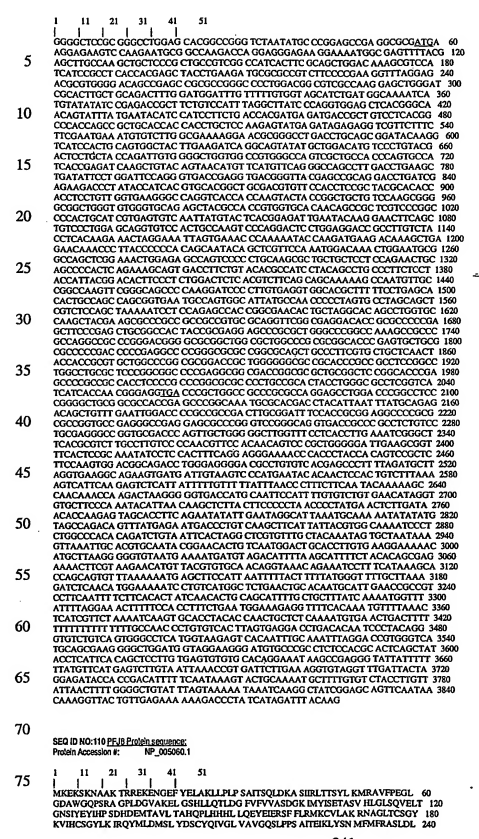
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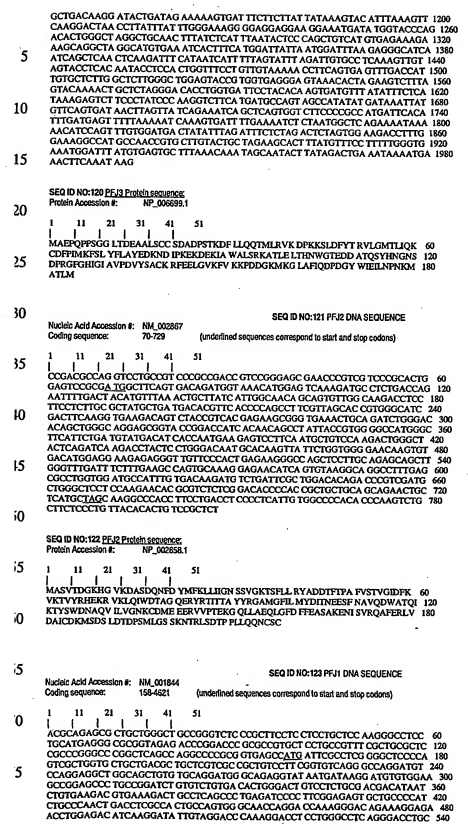
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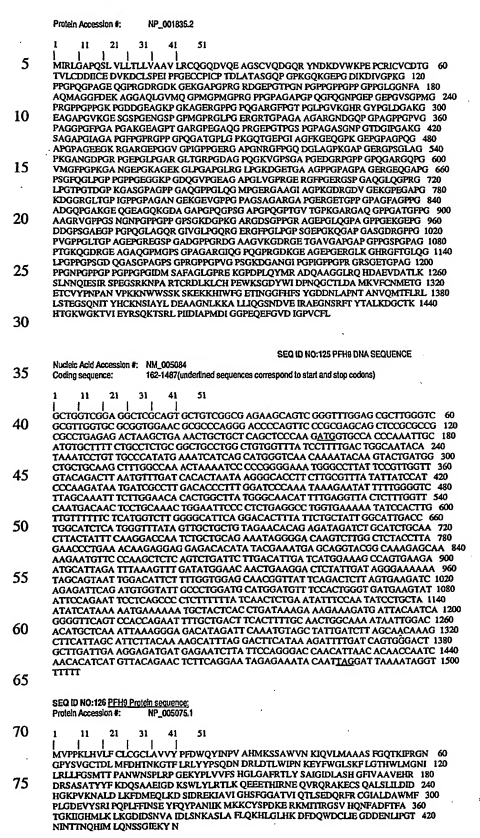
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TITTAGCTTT CCCATGCATA CTTAACTGCA CTTGCTTTAT CTCCTTGGGC ATTCGTACTT 1680 35 AGGATTCAAT AGAAACATGT ACAGGGTAAA CAATTTTTTA AAAATAAAAC TTCATGGAGT 1740 ΑΛΑΛΑΛΑ ΑΛΑΛΑΛΑ 40 SEQ ID NO:128 PFH8 Protein sequence Protein Accession #: NP 056984.1 45 51 11 21 31 41 MPPGPWESCF WVGGLILWLS VGSSGDAPPT POPKCADFOS ANLFEGTDLK VOFLLFVPSN 60 PSCGQLVEGS SDLQNSGFNA TLGTKLIIHG FRYLGTKPSW IDTFIRTILIR ATNANVIAVD 120 WIYGSTGVYF SAVKNVIKLS LEISLFLNKL LVLGVSESSI HIIGVSLGAH VGGMVGOLFG 180 50 GQLGQITGLD PAGPEYTRAS VEERLDAGDA LFVEAIHTDT DNLGIRIFVG HYDYFVNGGQ 240 DQPGCPTFFY AGYSYLICDH MRAVHLYISA LENSCPLMAF PCASYKAFLA GRCLDCFNPF 300 LLSCPRIGLY EQGGVKIEPL PKEVKVYLLT TSSAPYCMHH SLVEFHLKEL RNKDTNIEVT 360 FLSSNITSSS KITIPKQQRY GKGIIAHATP QCQINQVKFK FQSSNRVWKK DRTTIIGKFC 420 TALLPVNDRE KMVCLPEPVN LQASVTVSCD LKIACV 55 SEQ ID NO:129 PFH7 DNA SEQUENCE Nucleic Acid Accession #: NM_014384 60 Coding sequence: 89-1338 (underlined sequences correspond to start and stop codons) 21 31 41 65 CGTTGCCGGG TCGCAGGTCC CGCCAGTGCG AGCGCAACGG AGGTCGAAGG CGTTCAGACT 60 CTTIGCTGAA CGCGGAGCTG CGCCGGCTAT GCTGTGAGC GGCTGCAGGC GGTTCCGGGG 120
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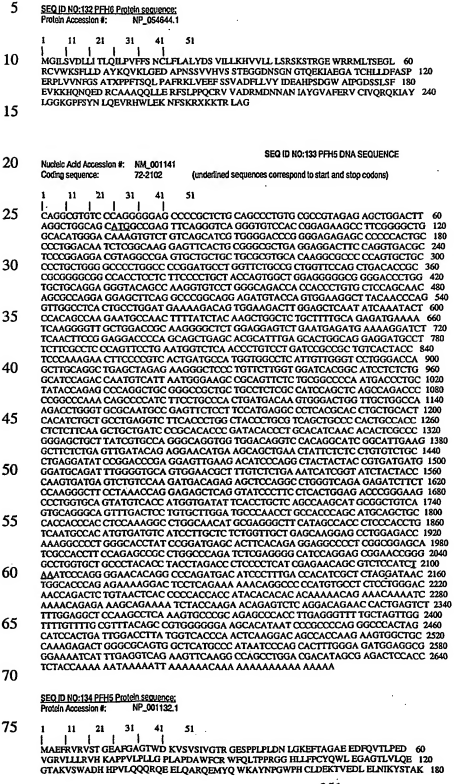
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SEQ ID NO:135 PFH4 DNA SEQUENCE

Nucleic Acid Accession #: NM_002742

Coding sequence: 236-2974 (underlined sequences correspond to start and stop codons)

1 11 21 31 41 5

5

10

15

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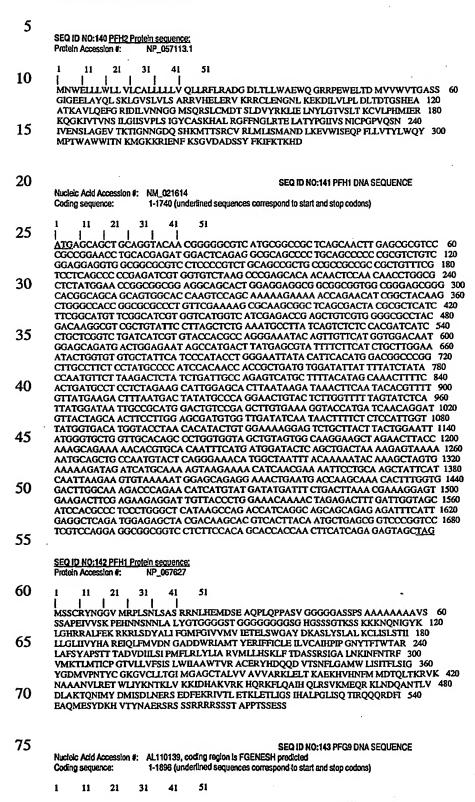
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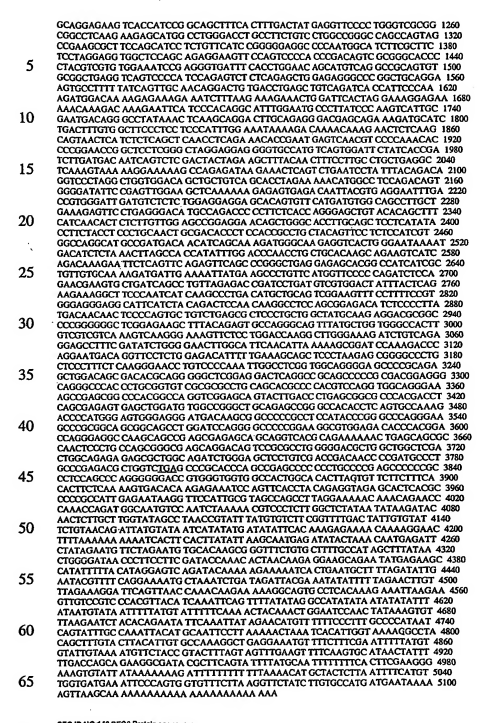
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CACCCCCCAGACCCC CCACCCTCGA CCCCCCTCGA CCCCCACACCGTC CCCGGACCCGA 1020 70 75 GAGGAAGCTG CGCCAGACCC GCAGCCTGGA CCCGGCCCTG ATCGGCGGCT GCGGGAGCGA 1020 CGAGGCGGGC GCGGAGGGCA GTGCGCGGGG AGCCACGGCG GGCCGCCTCT ACTCCCCATC 1080 ACTOCCAGOC GAGAGTOTOG GCCCTCGCTT GGCGTCCTCT TCCCGGGGTC CGCCCCCCAG 1140 GGCCACCAGG CTACCGCCTC CTGGACCTCT TTGCTCGTCC TTCTCCACAC CCAGCACCCC 1200



70 SEQ ID NO:146 PFG6 Protein sequence: Protein Accession #: NP_038286.1

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MSAQSILHSV FSCSSPASSS AASAKGFSKR KLRQTRSLDP ALIGGCGSDE AGAEGSARGA 60
TAGRLYSPSL PAESLGPRLA SSSRGPPPRA TRLPPPGPLC SSFSTPSTPQ EKSPSGSFHF 120
DYEVPLGRGG LKKSMAWDLP SVLAGPASSR SASSILCSSG GGPNGIFASP RRWLQQRKFQ 180
SPPDSRGHPY VVWKSEGDFT WNSMSGRSVR LRSVPIQSLS ELERARLQEV PFYQLQQDCD 240
LSCQITTPKD GQKRKKSLRK KLDSLGKEKN KDKEFTPQAF GMPLSQVIAN DRAYKLKQDL 300
QRDEQKDASD FVASLLPFGN KRQNKELSSS NSSLSSTSET PNESTSPNTP EPAPRARRRG 360

LTRELYTAFI NTLLLEPEEQ LGTLQLLIYL LPPCNCDTLH RLLQFLSIVA RHADDNISKD 540 GQEVTGNKMT SLNLATIFGP NLLHKQKSSD KEFSVQSSAR AEESTAIIAV VQKMIENYEA 600 5 LFMVPPDLQN EVLISLLETD PDVVDYLLRR KASQSSSPDM LQSEVSFSVG GRHSSTDSNK 660 ASSGDISPYD NNSPVLSERS LLAMQEDAAP GGSEKLYRVP GQFMLVGHLS SSKSRESSPG 720 PRLGKDLSEE PFDIWGTWHS TLKSGSKDPG MTGSSGDIFE SSSLRAGPCS LSQGNLSPNW 780 PRWQGSPAEL DSDTQGARRT QAAAPATEGR AHPAVSRACS TPHVQVAGKA ERPTARSEQY 840 LTLSGAHDLS ESELDVAGLQ SRATPQCQRP HGSGRDDKRP PPPYPGPGKP AAAAAWIQGP 900 10 PEGVETPTDQ GGQAAEREQQ VTQKKLSSAN SLPAGEQDSP RLGDAGWLDW QRERWQIWEL 960 LSTDNPDALP ETLY 15 **SEQ ID NO:147 PFG4 DNA SEQUENCE** Nucleic Acid Accession #: NM_002202 Coding sequence: 240-1289 (underlined sequences correspond to start and stop codons) 20 11 21 41 51 31 CCCCCGAGCC GCGCCGAGTC TGCCGCCGCC GCAGCGCCTC CGCTCCGCCA ACTCCGCCGG 60 CTTAAATTGG ACTCCTAGAT CCGCGAGGGC GCGGCGCAGC CGAGCAGCGG CTCTTTCAGC 120 ATTGGCAACC CCAGGGGCCA ATATTTCCCA CTTAGCCACA GCTCCAGCAT CCTCTCTGTG 180 25 GGCTGTTCAC CAACTGTACA ACCACCATTT CACTGTGGAC ATTACTCCCT CTTACAGATA 240 TGGGAGACAT GGGAGATCCA CCAAAAAAA AACGTCTGAT TTCCCTATGT GTTGGTTGCG 300 GCAATCAGAT TCACGATCAG TATATTCTGA GGGTTTCTCC GGATTTGGAA TGGCATGCGG 360 CATGITTGAA ATGTGCGGAG TGTAATCAGT ATTTGGACGA GAGCTGTACA TGCTTTGTTA 420 GGGATGGGAA AACCTACTGT AAAAGAGATT ATATCAGGTT GTACGGGATC AAATGCGCCA 480 30 AGTGCAGCAT CGGCTTCAGC AAGAACGACT TCGTGATGCG TGCCCGCTCC AAGGTGTATC 540 ACATCGAGTG TTTCCGCTGT GTGGCCTGCA GCCGCCAGCT CATCCCTGGG GACGAATTTG 600 CGCTTCGGGA GGACGGTCTC TTCTGCCGAG CAGACCACGA TGTGGTGGAG AGGGCCAGTC 660
TAGGCGCTGG CGACCCGCTC AGTCCCCTGC ATCCAGCGCG GCCACTGCAA ATGGCAGCGG 720
AGCCCATCTC CGCCAGGCAG CCAGCCCTGC GGCCCCACGT CCACAAGCAG CCGGAGAAGA 780
CCACCCGCGT GCGGACTGTG CTGAACGAGA AGCAGCTGCA CACCTTGCGG ACCTGCTACG 840 35 CCGCAAACCC GCGGCCAGAT GCGCTCATGA AGGAGCAACT GGTAGAGATG ACGGGCCTCA 900 GTCCCCGTGT GATCCGGGTC TGGTTTCAAA ACAAGCGGTG CAAGGACAAG AAGCGAAGCA 960 TCATGATGAA GCAACTCCAG CAGCAGCAGC CCAATGACAA AACTAATATC CAGGGGATGA 1020 CAGGAACTCC CATGGTGGCT GCCAGTCCAG AGAGACACGA CGGTGGCTTA CAGGCTAACC 1080 40 CAGTGGAAGT ACAAAGTTAC CAGCCACCTT GGAAAGTACT GAGCGACTTC GCCTTGCAGA 1140 GTGACATAGA TCAGCCTGCT TTTCAGCAAC TGGTCAATTT TTCAGAAGGA GGACCGGGCT 1200
CTAATTCCAC TGGCAGTGAA GTAGCATCAA TGTCCTCTCA ACTTCCAGAT ACACCTAACA 1260
GCATGGTAGC CAGTCCTATT GAGGCATCAA GTACCTCTCA TCTGTATTTT TTTTCCCTGT 1320
TGGAGAAAGT GGGAAATTAT AATGTCGAAC TCTGAAACAA AAGTATTTAA CGACCCAGTC 1380 45 AATGAAAACT GAATCAAGAA ATGAATGCTC CATGAAATGC ACGAAGTCTG TTTTAATGAC 1440 AAGGTGATAT GGTAGCAACA CTGTGAAGAC AATCATGGGA TTTTACTAGA ATTAAACAAC 1500 AAACAAAACG CAAAACCCAG TATATGCTAT TCAATGATCT TAGAAGTACT GAAAAAAAA 1560 GACGITITTA AAACGTAGAG GATITATATT CAAGGATCTC AAAGAAAGCA TITTCATITC 1620 ACTGCACATC TAGAGAAAAA CAAAAATAGA AAATTTTCTA GTCCATCCTA ATCTGAATGG 1680 50 TGCTGTTTCT ATATTGGTCA TTGCCTTGCC AAACAGGAGC TCCAGCAAAA GCGCAGGAAG 1740 AGAGACTGGC CTCCTTGGCT GAAAGAGTCC TTTCAGGAAG GTGGAGCTGC ATTGGTTTGA 1800 TATGTTTAAA GTTGACTTTA ACAAGGGGTT AATTGAAATC CTGGGTCTCT TGGCCTGTCC 1860 TGTAGCTGGT TTATTTTTA CTTTGCCCCC TCCCCACTTT TTTTGAGATC CATCCTTTAT 1920 CAAGAAGTCT GAAGCGACTA TAAAGGTTTT TGAATTCAGA TTTAAAAACC AACTTATAAA 1980

AMSVDSITDL DDNQSRLLEA LQLSLPAEAQ SKKEKARDKK LSLNPIYRQV PRLVDSCCQH 420 LEKHGLQTVG IFRVGSSKKR VRQLREEFDR GIDVSLEEEH SVHDVAALLK EFLRDMPDPL 480

65 SEQ ID NO:148 PFG4 Protein sequence: Protein Accession #: NP_002193.1

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70 MGDPPKKRL ISLCVGCGNQ IHDQYILRVS PDLEWHAACL KCAECNQYLD ESCTCFVRDG 60
KTYCKRDYIR LYGIKCAKCS IGFSKNDFVM RARSKVYHE CFRCVACSRQ LIPGDEFALR 120
EDGLFCRADH DVVERASLGA GDPLSPLHPA RPLQMAAEPI SARQPALRPH VHKQPEKTTR 180
VRTVLNEKQL HTLRTCYAAN PRPDALMKEQ LVEMTGLSPR VIRVWFQNKR CKDKKRSIMM 240
KQLQQQPND KTNIQGMTGT PMVAASPERH DGGLQANPVE VQSYQPPWKV LSDFALQSDI 300
DQPAFQQLVN FSEGGPGSNS TGSEVASMSS QLPDTPNSMV ASPIEA

GCATTGCAAC AAGGTTACCT CTATTTTGCC ACAAGCGTCT CGGGATTGTG TTTGACTTGT 2040 GTCTGTCCAA GAACTTTTCC CCCAAAGATG TGTATAGGTTA TTGGTTAAAA TGACTGTTTT 2100 CTCTCTCTAT GGAAATAAAA AGGAAAAAAA AAAGGAAACT TTTTTTGTTT GCTCTTGCAT 2160 TGCAAAAATT ATAAAGTAAT TTATTATTTA TTGTCGGAAG ACTTGCCACT TTTCATGTCA 2220 TTTGACATTT TTTGTTTGCT GAAGTGAAAA AAAAAGATAA AGGTTGTACG GTGGTCTTTG 2280

AATTATATGT CTAATTCTAT GTGTTTTGTC TTTTTCTTAA ATATTATGTG AAATCAAAGC 2340 GCCATATGTA GAATTATATC TTCAGGACTA TTTCACTAAT AAACATTTGG CATAGAT

SEQ ID NO:149 PFG2 DNA SEQUENCE

Nucleic Acid Accession #: NIM_001172 39-1103 (underlined sequences correspond to start and stop codons) Coding sequence: 5 GCGGAGCTCT GCCTTGGAGA TTCTCAGTGC TGCGGATCAT GTCCCTAAGG GGCAGCCTCT 60 CGCGTCTCCT CCAGACGCGA GTGCATTCCA TCCTGAAGAA ATCCGTCCAC TCCGTGGCTG 120 TGATAGGAGC CCCGTTCTCA CAAGGGCAGA AAAGAAAAGG AGTGGAGCAT GGTCCCGCTG 180 10 CCATAAGAGA AGCTGGCTTG ATGAAAAGGC TCTCCAGTTT GGGCTGCCAC CTAAAAGACT 240 TTGGAGATTT GAGTTTTACT CCAGTCCCCA AAGATGATCT CTACAACAAC CTGATAGTGA 300 ATCCACGCTC AGTGGGTCTT GCCAACCAGG AACTGGCTGA GGTGGTTAGC AGAGCTGTGT 360 CAGATGGCTA CAGCTGTGTC ACACTGGGAG GAGACCACAG CCTGGCAATC GGTACCATTA 420 GTGGCCATGC CCGACACTGC CCAGACCTTT GTGTTGTCTG GGTTGATGCC CATGCTGACA 480 15 TCAACACACC CCTTACCACT TCATCAGGAA ATCTCCATGG ACAGCCAGTT TCATTTCTCC 540 TCAACACACC CCTTACCACT TCATCAGGAA ATCTCCATGG ACAGCCAGTT TCATTTCTC 540
TCAGAGAACT ACAGGATAAG GTACCACAAC TCCCAGGATT TTCCTGGATC AAACCTTGTA 600
TCTCTTCTGC AAGTATTGTG TATATTGGTC TGAGAGACGT GGACCCTCCT GAACATTTTA 660
TTTTAAAGAA CTATGATATC CAGTATTTTT CCATGAGAGA TATTGATCGA CTTGGTATCC 720
AGAAGGTCAT GGAACGAACA TTTGATCTGC TGATTGGCAA GAGACAAAGA CCAATCCATT 780
TGAGTTTTGA TATTGATGCA TTTGACCCTA CACTGGCTCC AGCCACAGGA ACTCCTGTTG 840
TCGGGGGGACT AACCTATCGA GAAGGCATGT ATATTGCTGA GGAAATACAC AATACAGGGT 900 20 TGCTATCAGC ACTGGATCTT GTTGAAGTCA ATCCTCAGTT GGCCACCTCA GAGGAAGAGG 960 CGAAGACTAC AGCTAACCTG GCAGTAGATG TGATTGCTTC AAGCTTTGGT CAGACAAGAG 1020 AAGGAGGGCA TATTGTCTAT GACCAACTTC CTACTCCCAG TTCACCAGAT GAATCAGAAA 1080 25 ATCAAGCACG TGTGAGAATT TAGGAGACAC TGTGCACTGA CATGTTTCAC AACAGGCATT 1140 CCAGAATTAT GAGGCATTGA GGGGATAGAT GAATACTAAA TGGTTGTCTG GGTCAATACT 1200 GCCTTAATGA GAACATTTAC ACATTCTCAC AATTGTAAAG TTTCCCCTCT ATTTTGGTGA 1260 CCAATACTAC TGTAAATGTA TTTGGTTTTT TGCAGTTCAC AGGGTATTAA TATGCTACAG 1320 TACTATGTAA ATTTAAAGAA GTCATAAACA GCATTTATTA CCTTGGTATA TCATACTGGT 1380 CTIGITIGGTG TIGITICCTIC ACATITAAGT GGTTITICAT CTTTCCCCC TCCTCCCACA 1440
GCCTGGCTAT ACAGTGCATC CTTGAACTGT CAGCCCACAG CAGCAATATG CTTATTCTAT 1500
CCACATCCCT AACATCATGC ATTCACAAGG TCAAAGTTCT GGTCCACAAA CCCTTCCCTA 1560
TAGAAGTTCA ATGGCTGCGA AAGAATTTGT AGTAAACCAG GCCTCCCAGG ATGGCGAGCT 1620 30 CCAGTAAGAT GATAATGGAA AGCAGCAGCT TGTTGGTTGT CACTCTACAA AGAGAAGCAA 1680 35 AGTGGGGAGT AGTCAGAAGT TTGGATAACC TTCCTTCTAA ACATTTGGGG GTTAGACCTG 1740 GGACCACGGC TGGATACTCT GAGGCTGTAT GTTTGATCAC ACAGCCACTT AGCAGGAAGT 1800 ACTCATAAGG TTCTTTAGCT GTCACTTAGG GATAACACTG TCTACCTCAC AGAAATGTTA 1860 AACTGAGACA ATAAAACCCA AAGCAT 40 SEQ ID NO:150 PFG2 Protein sequence:
Protein Accession #: NP_001163.1 31 45 MSLRGSLSRL LQTRVHSILK KSVHSVAVIG APFSQGQKRK GVEHGPAAIR EAGLMKRLSS 60 LGCHLKDFGD LSFTPVPKDD LYNNLIVNPR SVGLANQELA EVVSRAVSDG YSCVTLGGDH 120 SLAIGTISGH ARHCPDLCVV WVDAHADINT PLTTSSGNLH GQPVSFLLRE LQDKVPQLPG 180 FSWIKPCISS ASIVYIGIRD VDPPEHFILK NYDIQYFSMR DIDRLGIQKV MERTFDLLIG 240 KRQRPIHLSF DIDAFDPTLA PATGTPVVGG LTYREGMYIA EEIHNTGLLS ALDLVEVNPQ 300 LATSEEEAKT TANLAVDVIA SSFGQTREGG HIVYDQLPTP SSPDESENQA RVRI 50 55 SEQ ID NO:151 PFG1 DNA SEQUENCE Nucleic Acid Accession #: NM_017906 80-1255 (underlined sequences correspond to start and stop codons) Coding sequence: 51 60 AATTATATAT TITTACTCTA TGTTTCTCTA CATGTTTTTT TCTTTCCGTT GCTGGCGGAA 60
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GTTCGCTGTA CACCCGGAGC CCAAGGCTTG CGGCGACCAC GAGCAATGGA CTCTTGTGGC 180
TGACTTCACT CACCATGCTC ACACTGCCTC CTTGTCAGCA GTAGCTGTAA ATAGTCGTTT 240
TGTGGTCACT GGGAGCAAGA ATGAAACAAT TCACATTTAT GACATGAAAA AGAAGATTGA 300
GCATGGGGCT CTAGTGCATC ACAGTGGTAC AATAACTTGC CTGAAAATTCT ATGGCAACAG 360 65 GCATTTAATC AGTGGAGCGG AAGATGGACT CATCTGTATC TGGGATGCAA AGAAATGGGA 420 ATGCCTGAAG TCAATTAAAG CTCACAAAGG ACAGGTGACC TTCCTTTCTA TTCACCCATC 480 TGGCAAGTTG GCCCTGTCGG TTGGTACAGA TAAAACTTTA AGAACGTGGA ATCTTGTAGA 540 70 AGGAAGATCA GCATTCATAA AAAATATAAA ACAAAATGCT CACATAGTAG AATGGTCCCC 600 AAGAGGAGAG CAGTATGTAG TTATCATACA GAATAAAATA GACATCTATC AGCTTGACAC 660 TGCATCCATT AGTGGCACCA TCACAAATGA AAAGAGAATT TCCTCTGTTA AATTTCTTTC 720 AGAGTCTGTC CTTGCAGTGG CTGGAGATGA AGAAGTTATA AGGTTTTTTG ACTGTGATTC 780 ACTAGTGTGC CTCTGCGAAT TTAAAGCTCA TGAAAACAGG GTAAAGGACA TGTTCAGTTT 840 75 TGAAATTCCA GAGCATCATG TTATTGTTTC AGCATCGAGT GATGGTTTCA TCAAAATGTG 900
GAAGCTTAAG CAGGATAAGA AAGTTCCCCC ATCTTTACTC TGTGAAATAA ACACTAATGC 960
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AGGTGACAGT AAGAAAGCAA CAAAAGAAAG TGGCCTGATA TCAACCAAGA AGAGGAAAAT 1200 GGTAGAAATG TTGGAAAAGA AGAGGAAAAA GAAGAAAATA AAAACAATGC AG<u>TGA</u>ATCAC 1260 AGATGTCTCC TGAAAGAACT CTTTTAGATG AAATCATTCT ACTCAAATGT ACCTTAATTT 1320
TTTTTTTTCC CTGAGTAAAA GCAAGAAATT TCTTCCTTTG GAAAAAATAT ATATATTAAA 1380
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10 SEQ ID NO:152 PFG1 Protein sequence: Protein Accession #: NP_060376.1

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31

15 MELVAGCYEQ VLFGFAVHPE PKACGDHEQW TLVADFTHHA HTASLSAVAV NSRFVVTGSK 60 DETHHYDMK KKIEHGALVH HSGTITCLKF YGNRHLISGA EDGLICIWDA KKWECLKSIK 120
AHKGQVTFLS IHPSGKLALS VGTDKTLRTW NLVEGRSAFI KNIKQNAHIV EWSPRGEQYV 180
VIIQNKIDIY QLDTASISGT ITNEKRISSV KFLSESVLAV AGDEEVIRFF DCDSLVCLCE 240
FKAHENRVKD MFSFEIPEHH VIVSASSDGF IKMWKLKQDK KVPPSLLCEI NTNARLTCLG 300 20 VWLDKVADMK SLPPAAEPSP VSKEQSKIGK KEPGDTVHKE EKRSKPNTKK RGLTGDSKKA 360 TKESGLISTK KRKMVEMLEK KRKKKKIKTM Q

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SEQ ID NO:153 PFD6 DNA SEQUENCE

Nucleic Acid Accession #: NM_014668 (underlined sequences correspond to start and stop codons) Coding sequence: 110-2953

51

GATGTCTTGG ACATGCTCTG GCTGGCTAAT CTCCATGTTC TAGCCGACTG AAAATACGGT 60 GGCCAAGTGG ATGGTGTGCT TATTTGCAGT CTAAAGAAAT TTCCTTTTGA TGTGGCAGAA 120 AATCGAGGAT GTGGAGTGGA GACCCCAGAC TTACTTGGAG CTGGAGGGTC TGCCTTGCAT 180 35 CCTGATCTTC AGTGGGATGG ACCCGCATGG GGAGTCCTTG CCGAGGTCTT TGAGGTACTG 240

TGACCTGCGA TTGATAAACT CCTCCTGCTT GGTGAGAACA GCCTTGGAGC AGGAGCTGGG 300
CCTGGCTGCC TACTTTGTGA GCAACGAGGT TCCCTTGGAG AAGGGGGGCTA GGAACGAGGC 360
CTTGGAGAGT GATGCTGAGA AGCTGAGCAG CACAGACAAC GAGGATGAGG AGCTGGGGAC 420
AGAAGGCTCT ACCTCGGAGA AGAGAAGCCC CATGAAAAGG GAGAGGTCCC GCTCCCACGA 480 40 CTCAGCATCC TCATCCCTCT CCTCCAAGGC TTCCGGTTCA GCGCTCGGTG GCGAGTCCTC 540 GGCTCAGCCC ACAGCACTCC CCCAGGGAGA GCATGCCAGG TCGCCCCAGC CCCGTGGCCC 600 CGCAGAGGAG GGCAGAGCCC CTGGTGAGAA ACAGAGGCCC CGGGCAAGTC AGGGGCCACC 660

CTCGGCCATC AGCAGGCACA GTCCCGGGCC GACGCCCCAG CCCGACTGTA GCCTCAGGAC 720 CGGCCAGAGG AGCGTCCAGG TGTCGGTCAC CTCGTCGTGC TCCCAGCTGT CCTCCTCCTC 780 45 GGGCTCATCC TCCTCATCCG TGGCGCCCGC TGCCGGCACG TGGGTCCTGC AGGCCTCCCA 840 GTGCTCCTTG ACCAAGGCCT GCCGCCAGCC ACCCATTGTC TTCTTGCCCA AGCTCGTGTA 900

CGACATGGTT GTGTCCACTG ACAGCAGTGG CCTGCCCAAG GCCGCCTCCC TCCTGCCCTC 960 CCCCTCGGTC ATGTGGGCCA GCTCTTTCCG CCCCTGCTC AGCAAGACCA TGACATCCAC 1020 CGAGCAGTCC CTCTACTACC GGCAGTGGAC GGTGCCCCGG CCCAGCCACA TGGACTACGG 1080 CAACOGGGCC GAGGGCCGCG TGGACGGCTT CCACCCCCGC AGGCTGCTGC TCAGCGGCCC 1140 CCCTCAGATC GGGAAGACAG GTGCCTACCT GCAGTTCCTC AGTGTCCTGT CCAGGATGCT 1200 TGTTCGGCTC ACAGAAGTGG ATGTCTATGA CGAGGAGGAG ATCAATATCA ACCTCAGAGA 1260 AGAATCTGAC TGGCATTATC TCCAGCTTAG CGACCCCTGG CCAGACCTGG AGCTGTTCAA 1320

GAAGTTGCCC TTTGACTACA TCATTCACGA CCCGAAGTAT GAAGATGCCA GCCTGATTTG 1380 55

60 GGAGAGCATG CGACTACCCC TCGTGACAGA CAAGAGCCAT GAATATATAA AAAGTCCGAC 1740 ATTCACTCCA ACCACOGGCC GTCACGAACA TGGGCTCTTT AATCTGTACC ACGCAATGGA 1800 CGGTGCCAGC CATTTGCACG TGCTGGTTGT CAAGGAATAC GAGATGGCAA TTTATAAGAA 1860 ATATTGGCCC AACCACATCA TGCTGGTGCT CCCCAGTATC TTCAACAGTG CTGGAGTTGG 1920

TGCTGCTCAT TTCCTCATCA AGGAGCTGTC CTACCATAAC CTGGAGCTCG AGCGGAACCG 1980 GCAGGAGGAG CTGGGAATCA AGCCGCAGGA CATCTGGCCT TTCATTGTGA TCTCTGATGA 2040 CTCCTGCGTG ATGTGGAACG TGGTGGATGT CAACTCTGCT GGGGAGAAA GCAGGGAGTT 2100 CTCCTGGTCG GAAAGGAACG TGTCTTTGAA GCACATCATG CAGCACATCG AGGCGGCCCC 2160 CGACATCATG CACTACGCCC TGCTGGGCCT GCGGAAGTGG TCCAGCAAGA CCCGGGCCAG 2220 CGAGGTGCAA GAGCCCTTCT CCCGCTGCCA CGTGCACAAC TTCATCATCC TGAACGTGGA 2280 65

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CGCAGCGCCC GCCCAGCTCC TGCTGGAGAA GTTCCTGCAG CACCACAGCC ACCTCTTCTT 2880 CCCGCTGTCC CTGAAGAACC ATGACCACCC AGTGCTGTCT GTCGACTGTT ACCTGAACCT 2640 75 GGGATCTCAG ATTTCTGTTT GCTATGTGAG CTCCAGGCCC CACTCTTTAA ACATCAGCTG 2700 CTCGGACTTG CTGTTCAGTG GGCTGCTGCT GTACCTCTGT GACTCTTTTG TGGGAGCTAG 2760 CTTTTTGAAA AAGTTTCATT TTCTGAAAGG TGCGACGTTG TGTGTCATCT GTCAGGACCG 2820 GAGCTCACTG CGCCAGACGG TCGTCCGCCT GGAGCTCGAG GACGAGTGGC AGTTCCGGCT 2880

GCGCGATGAG TTCCAGACCG CCAATGCCAG GGAAGACCGG CCGCTCTTTT TTCTGACGGG 2940 ACGACACATC TGAGGAAGAC AGCGGCGAGT TTTCTGAAGA GATGAGTGCT CAGAGCCCTC 3000 ATGCTGTTGA GGCTAAAGGG AGGCCTGGAA CGGTGGGGGCG TTTGACTGGA ATGGACCCCA 3060 GGGACTGTCC AGGTGCAGCC CCTCCTAGTA CACATGGGCC CCCGAGGCCG TGGTCCTGGG 3120 5 AGCCAGGAAG ACTCCGCAGT GGGTGAGAAT GAAAACTTGA GACTCCCAAG TTCTGGGCCA 3180 GCCCATTGCT CTGGGCTGTT TTAAAGCCCA TTTCACGAGG AACAAAGATT TACTTCCTGT 3240 CCTGCCATTC GTGTGCTTCC ATGGACAAAC CTGATTTTTT TCTCTTAGTT CTAAAGAATC 3300 TTGGGTTATT TTGTAGCGGT GCCAGTATTT CAGTAGATGG GATTTCAGCC AAGTAGGTTC 3360 CCCTGTAACC TCCTACAAAG CAATATTCCA AAGGAACATT TTAACTGTAA AGGCTGGAGA 3420 10 CAAGAAAAA TAAGTAGATC GTTTTAATAA CAATTATTTA ATTGCCTATA AGTTTGCTGT 3480 TTCAGAGGCT AGCCCAAAGG CATCAAATTT AATAAAGTTA AACAAATTGA TTTACTTCAG 3540
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CATTGCCCGG CGCCTGGACA GTGATCATCT TGTTCATCTT GTTCAGTCCT TTCTTGTGTG 4380 25 ATTGGAATTA TTCATCCCCT TTGAAAGATG AGAAGGTTGA GATGCAAAGA GTCTACCTTT 4440 CCAAGTTCTC ACTGCTGGAA AGAGCTAGAA GCACAGTTCA AAGTTCTGGC TTCTGGACTC 4500 TGCAGTCCAG GTCTCCCTTC TCCCACTTGC CTACCCTCAA TGCCACACTG TTTTTGAAGT 4560 GGCCCATAAC TTGAAGGAAA AGTTTAAAGA CAGTTCAATT TAATCATCAG AATGCATTCT 4620 TTTTTTTTC GGAGGGAG TTTCACTCTT GCTGCCAGG CTGGAGTGCA ATGGCAT1CT 4620
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45 SEQ ID NO:154 PFD6 Protein sequence: Protein Accession #: NP_055483.1

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1 11 21 31 41 51

50 MWQKIEDVEW RPQTYLELEG LPCILIFSGM DPHGESLPRS LRYCDLRLIN SSCLVRTALE 60
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RSHDSASSSL SSKASGSALG GESSAQPTAL PQGEHARSPQ PRGPAEEGRA PGEKQRPRAS 180
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MTSTEQSLYY RQWTVPRPSH MDYGNRAEGR VDGFHPRRLL LSGPPQIGKT GAYLQFLSVL 360
SRMLVRLTEV DVYDEFEINI NLREESDWHY LQLSDPWPDL ELFKKLPFDY IIHDPKYEDA 420
SLICSHYQGI KSEDRGMSRK PEDLYVRRQT ARMRLSKYAA YNTYHHCEQC HQYMGFHPRY 480
QLYESTLHAF AFSYSMLGEE IQLHFIIPKS KEHHFVFSQP GGQLESMRLP LVTDKSHEYI 540

SLICSHYQGI KSEDRGMSRK PEDLYVRRQT ARMRISKYAA YNTYHHCEQC HQYMGFHPRY 480 QLYSTLHAF AFSYSMLGEE IQLHFIIPKS KEHHFVFSQP GGQLESMRLP LVTDKSHEYI 540 KSPIFIPTTG RHEHGLFNLY HAMDGASHLH VLVVKEYEMA IYKKYWPNHI MLVLPSIFNS 600 AGVGAAHFLI KELSYHNLEL ERNRQEELGI KPQDIWPFIV ISDDSCVMWN VVDVNSAGER 660 SREFSWSERN VSLKHINQHI EAAPDIMHYA LLGLRKWSSK TRASEVQEPF SRCHVHNFII 720 LNVDLTQNVQ YNQNRFLCDD VDFNLRVHSA GLLLCFRNRF SVMKKQIVK GHRSFHITSK 780 VSDNSAAVVP AQVICAPDSK HTFLAAPAQL LLEKFLQHHS HLFFPLSLKN HDHPVLSVDC 840 YLNLGSQISV CYVSSRPHSL NISCSDLLFS GLLLYLCDSF VGASFLKKFH FLKGATLCVI 900

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SEQ ID NO:155 PFC6 DNA SEQUENCE

70 Nucleic Acid Accession *: NM_000522 Coding sequence: 1-1167 (underlined sequences correspond to start and stop codons)

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75
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GCCGCGTCCG CCTACAGCAG CGCCCCCGGG GAGGCGCCCC CGTCGGCTGC CGCCGCTGCT 360

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GCCGCGGCTG CCGCTGCAGC CGCCGCCGCCGCCGCGT CGTCCTCGGG AGGTCCCGGC 420 CCGGCGGGCC CGGCGGCGGC AGAGGCGGCC AAGCAATGCA GCCCCTGCTC GGCAGCGGCG 480 CAGAGCTCGT CGGGGCCCGC GGCGCTGCCC TATGGCTACT TCGGCAGCGG CTACTACCCG 540 CAGAGCILUIT CGGGGCCCGC GGCGCTGCCC TATGGCTACT TCGGCAGCGG CTACTACCCG 540
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AGGGGGAGAA AGAAGCGCCT GCCTTATACC AAGGTGCAAT TAAAAGAACT TGAACGGGAA 1020 10 TACGCCACGA ATAAATTCAT TACTAAGGAC AAACGGAGGC GGATATCAGC CACGACGAAT 1080 CTCTCTGAGC GGCAGGTCAC AATCTGGTTC CAGAACAGGA GGGTTAAAGA GAAAAAAGTC 1140 ATCAACAAAC TGAAAACCAC TAGT<u>TAA</u> 15 SEQ ID NO:156 PFC6 Protein sequence: Protein Accession #: NP_000513.1 20 11 31 41 51 21 MTASVLLHPR WIEPTVMFLY DNGGGLVADE LNKNMEGAAA AAAAAAAAA AGAGGGGFPH 60 PAAAAAGGNF SVAAAAAAA AAAANQCRNL MAHPAPLAPG AASAYSSAPG EAPPSAAAAA 120 AAAAAAAA AAASSSGGPG PAGPAAAEAA KQCSPCSAAA QSSSGPAALP YGYFGSGYYP 180 CARMGPPPNA IKSCPQPPSA AAAAAFADKY MDTAGPAAEE FSSRAKEFAF YHQGYAAGPY 240 25 HIHOPMPGYL DMPVVPGLGG PGESRHEPLG LPMESYQPWA LPNGWNGOMY CPKEQAQPPH 300 LWKSTLPDVV SHPSDASSYR RGRKKRVPYT KVQLKELERE YATNKFITKD KRRRISATTN 360 LSERQVTIWF QNRRVKEKKV INKLKTTS 30 SEQ ID NO:157 PFA3 DNA SEQUENCE Nucleic Acid Accession #: AW102723 523-2676 (underlined sequences correspond to start and stop codons) Coding sequence: 35 51 CCCTTATGGC GATTGGGCGG CTGCAGAGAC CAGGACTCAG TTCCCCTGCC CTAGTCTGAG 60 CCTAGTGGGT GGGACTCAGC TCAGAGTCAG TTTTCAGAAG CAGGTTTCAG TTGCAGAGTT 120 40 TTCCTACACT TTTCCTGCGC TAGAGCAGCG AGCAGCCTGG AACAGACCCA GGCGGAGGAC 180
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NGIRRIMNRR DFQKFMPNEY FEILTPKINQ TFSGIMTMLN MQFVVRVRRW DNSVKKSSRV 360
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51

SEQ ID NO:159 PFA1 DNA SEQUENCE

Nucleic Acid Accession #: NM_004362
Coding sequence: NM_004362
102-1934 (underlined sequences correspond to start and stop codons)

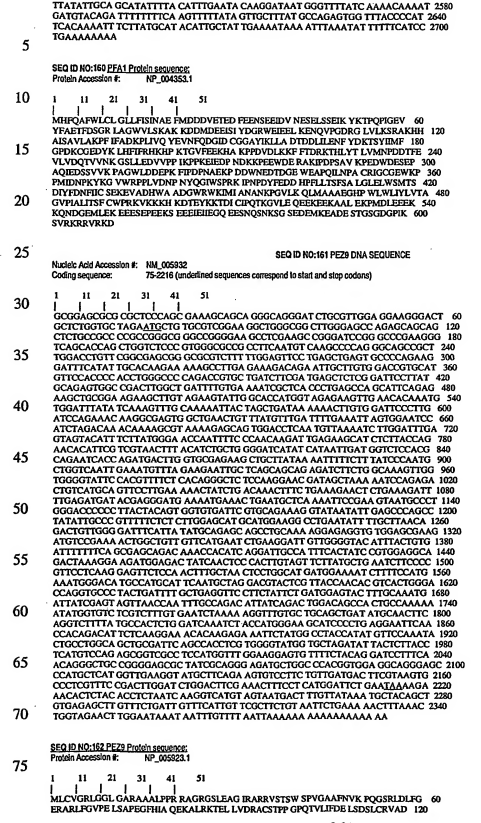
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SEQ ID NO:163 PEZ8 DNA SEQUENCE

Nucleic Acid Accession #: AF103907
Coding sequence: none (underlined sequences correspond to start and stop codons)

15

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PEZ8 Protein sequence:

10 Protein Accession #: none SEQ ID NO:164 PEZ6 DNA SEQUENCE Nucleic Acid Accession #: AB028945 1-3765 (underlined sequences correspond to start and stop codons) Coding sequence: 15 ATGATGATGA ACGTCCCCGG CGGAGGAGCG GCCGCGGTGA TGATGACGGG CTACAATAAT 60 GGTCGCTGTC CCCGGAATTC TCTCTACAGT GACTGCATTA TTGAGGAGAA GACGGTGGTC 120 CTGCAGAAAA AAGACAATGA GGGCTTTGGA TTCGTGCTTC GAGGGGCCAA AGCTGACACA 180 CCCATTGAAG AATTCACACC AACACCGGCT TTCCCAGCCC TACAGTACCT GGAGTCCGTG 240 20 GATGAAGGTG GGGTGGCGTG GCAAGCCGGA CTAAGGACCG GGGACTTCTT GATTGAGGTT 300 AACAATGAGA ATGTTGTCAA AGTCGGCCAC AGGCAGGTGG TGAACATGAT CCGGCAGGGA 360 GGGAATCACC TGGTCCTTAA GGTGGTCACG GTGACCAGGA ATCTGGACCC CGACGACACC 420 25 GCCAGGAAGA AAGCTCCCCC GCCTCCAAAG CGGGCACCGA CCACAGCCCT CACCCTGCGC 480 TCCAAGTCCA TGACCTCGGA GCTGGAGGAG CTCGTGGATA AAGATAAACC CGAGGAGATA 540 GTCCCGGCCT CCAAGCCCTC CCGCGCTGCT GAGAACATGG CTGTGGAACC GAGGGTGGCG 600 ACCATCAAGC AGCGGCCCAG CAGCCGGTGC TTCCCGGCGG GCTCAGACAT GAACTCTGTG 660 TACGAACGCC AAGGAATCGC COTGATGACG CCCACTGTTC CTGGGAGCCC AAAAGCCCCG 720
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ΑΛΑΓΑΛΑΛΑ ΑΛΑΛΑΛΑΛΑ ΑΛΑΛΑΛΑΛ

5 SEQ ID NO:165 PEZ6 Protein sequence; Protein Accession #; BAA82974.1

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SEQ ID NO:166 PEZ4 DNA SEQUENCE

Nucleic Acid Accession #: NM_000024
Coding sequence: NM_000024
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PCT/US01/32045 **WO** 02/30268

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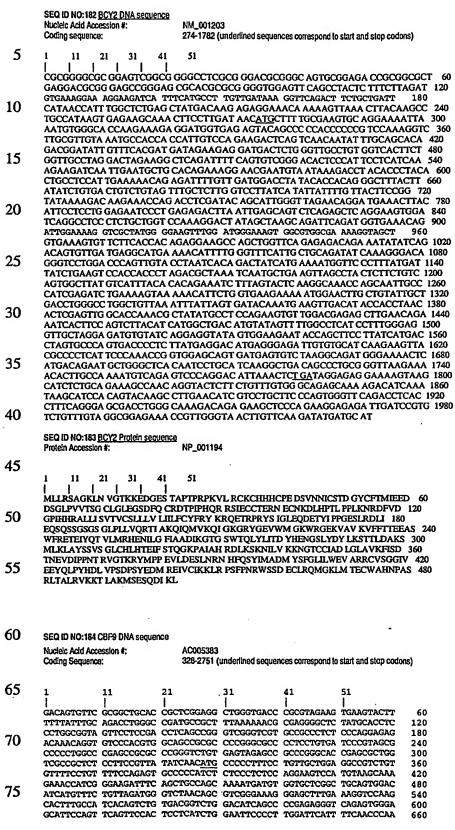
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50	1 GTCATATTGA CTTTGAACTC CGGAAAACCC CTCAGTACTA ACCCCGTCGT AGAAAGCACT CTGGCCTACT CTGGCCTACT CAGGAGGACGA GAGGACGA GAGGACGA ATACAAGCG CATCACTGGG CGCTCCCGGG GAGGCTCCAT TTAACAATC ATGACACCC ATGAGACCC ATGAGACCCGG	11 ACATTCCAGA AGGGTCACCA CTATCCCGCA CCGTCCCCC CTGCACGCAG GTGCATCAAC CTGCATCAAC CAGATCGTGT GAAGTCCTGG CAGGGACATG ATCCACCAGT GGTCAACTTG GGCCTGGCCC CATCACCTC CATGCCATTGG ATACCACCAGT ATACCACCAGT ATACCACAGT ATACCACCAGT ATACCACAGT ATACCACAGT ATACCACAGT ATACCACAGT ATACCACAGT ATACCACAGT ATACCACAGT	21 TACCTATCAT CCAGCTATTG CAGCCCACTG GTGCCCCAGT CCCAAATCCC TTGACCCTGG ATGGCCACCA CCCTCTAACT CACCCTGTGT GGCTATAAGA TTTATGAAAC GATGCCTGTT AACTCAAGCC TGGCAGGTCA ACGCAGGTCA ACGCAGTTTG CAAAAAGTCA	31 TACTCGATGC GACCTTACTA TGGTCCCCAC ACGCCCCAGA GGACCTTCCT AGTGCTCCAA GCCAAGACGAA ACCAAGACGA ATAATTTTA TGAACACAGA CCCAGAGCAG GCCAGAGCAG GCCAGAGCAG GCCAGAGCAG GCCTGCACGT TGACAGCCCG TGACAGCCCG TGACAGCCCG TGACAGCCCG	41 TGTTGATAAC TGAAAACCAT TGTCTACGAG GGTCCTGACG CGTGGGAGCT CTCTGGGATA CGTGTCACAC CTTCATCCTT CTGGAACGAG CTCTAGCCAA TGCCGGCAAT TGCTGGGACGCAC GATCGTGGGC CCACACGCTC CCACACCCCT AAATTATGAC	51 AGCAAGATEG GGATACCAAC GTGCATCCGG CAGGCTTCCA TCAAGACTA GCGCTGGCCG GAGTGCGACT TGCCCGGCG CAGATGTACT AACTACGGGC GGAATAGTGG GTCGATATCT TTACCGTGTT GGTGAGACCG CACGTGTGCG CACGTGTCCG CACATGTTCT TCACACACCA	60 120 180 240 300 360 420 480 540 660 720 780 840 900 960 1020 1080
50 55	1 GTCATATTGA CTTTGAACTC CGGARAACCC CTCAGTACTA ACCCGTCGT AGAAAGCACT CCTCAGGTAC CGTCAGGTAC GGGCGGCCTG ATGACAGCG ATAAAAAACT TAGCCTGCGG GAGGCTCCAT TTAACAATCC ATGACCGG ATGACCAGCGG GAGGCTCCAT TTAACAATCC ATGACCAGCGG AGACCAATGA	11 ACATTCCAGA AGGGTCACCA CCGTCCCC CTGCACGCA GRAGTTC CTGGAAGTTC CTGCATCAAC GAATCGGTGT CAGGGACATG ATCACCAGG GTACCACGC CTACCACGC CTACCACCAGC CTACCACCAGC CTACCACTG CGTCACTTG GGCCTGGCCC CATCACCCC ATGGCATTGG CATCACGTAC CATTGGCCTG CATTGGCCTG	21 TACCTATCAT CAGCCACTG CAGCCACTG GTGCCCAGT TTGACCCTGG ATGGGCAGCA CCCTCTAACT CACCCTGTGT GGCTATAAGA TTTATGAAAC TATGACTGG TTGACTGT GACTGAGTCA GATGCAGTCA GATGGATTG ACGCAGTTCA CAGGCATTTG CAAAAAGTGA ATGAAAGTGA ATGAAAGTGA	31 TACTCGATGC GACCTTACTA ACGCCCCAGA GCCCCCCAGA GGTGCTCCTA AGTGCTCCAA GGTGTGATGG ACGGACCAA ATAATTTTA TGAACACAAG GCCAGAGCAG GCCAGAGCAG GCCTGCACGT TGACAGCCGC CGGGGATTTT TTTCTCATCC AGAAGCCTCT	41 TGTTGATAAC TGAAAACCAT TGTCTACGAG GGTCCTGACG AGTGTGCACC CGTGGGAGCT CTCTGGGATA CGTGTCACCAC CTCTACCCAA CGTGCACAC CTCTAGCCAA TGCCGGCAAT AGTGGTTTCT CCACTGCGGC CCAGAACGTC CCACTGCGTG GAGACAAT CACTGCGTG GAGACAAT CACTGCGTG GAGACAAT CACTGCGTG GAGACAAT CACTGCAC GACATTCAAC GACTTTCAAC	51 AGCAAGATEG GGATACCAAC GTGCATCCGG CAGGCTTCCA TCAAAGACTA GGCTGGCCG GAGTGCGACT TGCCCGGCG CAGATGTACT TACCGGCC GGAATATCT TTACGCTGTT TTACGCTGTT TTACGCTGTT TCACAGACCC GAAAAACCTC TTCATGTTCT TCCAAGACCA GACCTAGTCA	60 120 180 240 300 420 480 540 660 720 780 900 960 1020 1080 1140
50 55 60	1 GTCATATTGA CTTTGAACTC CGGAAAACCC CTCAGTACTA ACCCGTCGT CTGACCTACT CCTCAGGTACT CCTCAGGTACT CCTCAGGTACT CATCAGGACACT ATGACACT ATGACACGG GAGGCTCCAT TAACAATCC ATGAGCCCA ATAAAAACC TTAACAATCC ATGAGCCGG AGACAATGA AACCAGTGTG AACCAGTGTG	11 ACATTCCAGA AGGGTCACCA CCGTCCCCC CTGCACGCAG GTGCATCACC CTGGATCACC CTGCATCACC GAATCGGTGT GAATCGGTGT GAAGTCCTGG ATCCACCAGC GTACCACAGT GTACCACAGT GTACCACAGT ATCCACCAGC CATCACCTC CATCACCTC ATGCCATTGG ATACCAACTT CATTGCGCTTG TCTGCCCAAC	21 TACCTATCAT CCAGCTATTG CAGCCACTG GTGCCCCAGT CCCAAATCCC TTGACCCTGG ATGGCAGCT CACCTGTATA GACCTGTT AACTCAAGCC TGTATAGAAC GATGCAGGTC AGCGAGGTC AACTCAAGCC AGGGAGTTG AACTCAAGCC CCAGGCATTG CAAAAAGTGA ATGAAGCTGC CCAGGCATGA	31 TACTCGATGC GACCTTACTA TGGTCCCCAC ACGCCCCAGA GGACCTTCCT AGTGCTCCAA GCCAAGACGAA ACCAAGACGA ATAATTTTA TGAACACAGA CCCAGAGCAG GCCAGAGCAG GCCAGAGCAG GCCAGAGCAG GCCTGCACGT TGACAGCCCG TGACAGCCCG TGACAGCCCG TGACAGCCCG	41 TGTTGATAAC TGAAAACCAT TGTCTACGAG GGTCCTGACG AGTGTGCACC CCTCGGGACC CCTCATCCTT CTGGAACGAG CTCTAGCCAA AGTGGTTCAT GATCGTGGCAA AGTGGTTTCT GATCGTGGGC CCACTGCGTG GAGACATCT AAATTATGAC AGAACACTC AGAACACTC AAATTATCAAC AGAACACTC	51 AGCAAGATGG GGATACCAAC GTGCATCCGA CAGGCTTCCA CAGGCTGCCC GAGTGCCACT AACTACGGGC CAGATGTACT AACTACGGGC GGAATACTGG GTCGATACT TTACGCTGTT GGTGAGACCA CAAGACCA TCCAAGACCA TCCAAGACCA TCCTAGTGA TCCTAGTGA TCCTAGTGA TCCTAGTGA TCCTAGTGA TCCTAGTGA TCCTAGTGA TCCTAGTGATT TCCAAGACCA TCCTAGTGA TCCTAGTGATT	60 120 180 240 300 360 420 480 540 660 720 780 840 900 960 1020 1080
50 55	1 GTCATATTGA CTTTGAACTC CGGAAAACCC CTCAGTACTA ACCCCGTCGT AGAAAGCACT CTGGCCTACT CTGGCCTACT CAGGAGACGA GGGCGGCCTG ATGACAGGG ATAAAAACT TTAACAATCC ATGAGGCCG AGACCACGG AGAACAATGA AACCAGTGTG CCGGGTGGGG TCCATGGGGGGTGCCAT CATGGAGCCGG AGAACAATGA AACCAGTGTG CCGGGTGGGG TGCTTCTCAT	11 ACATTCCAGA AGGGTCACCA CTATCCCGCA CCGTCCCCC CTGCACGCAG GTGCATCACC CTGCATCACC CTGCATCACC CAGGAGTTC CAGGAGTTC CAGGACATG ATCCACCAGT GGTCACACTG GGCACTGGCC CATCACCTC CATGCATTG ATACCACAGT GGCATTGG ATACCACAGT CATTGCCACAGT CATTGCCCAC CTCCCCAC CATGCCACCAGT CATGCCACCAGT CATGCCACCAGT CATGCCCACC CTCCCCACC CTCCCCCAC CTCCCCCACC CTCCCCCACC CTCCCCACC CTCCCCCC CTCCCCCC CTCCCCCCC CTCCCCCCC	21 TACCTATCAT CAGCCACTG CAGCCACTG GTGCCCAGT CCCAAATCCC TTGACCTGG ATGGCAGCA CCCTCTAACT GTTCGCCTCT GTTCGCCTCT GTTATGAAAC GATGCCTGTT AACTCAAGCC TGGCAGGTCA ACGCAGTCA ACGCAGTCA ACGCAGTCA ACGCAGTCA ACGCAGTCA ACGAAAAGTCA ATGAAACTCA ATGAAACTCA ATGAAAAGTCA ATGAAAAGTCA AGAGAAAAGGA AGAGAAAAGGA AGAGAAACGAA	31 TACTCGATGC GACCTTACTA TGGTCCCCAC ACGCCCCAGA GCACTTCCT AGTGCTCCAA GCCAAGACGA ACAGACGA ATAATTTTA TGAACACAGA GCCAGAGCCAG GCCAGAGCCGC CCGCGGATTT TTCTCATCC AGAAGCCTCT AGAAGCCTCT AGAAGCCTCA AGACCTCAGA GCAGATATT	41 TGTTGATAAC TGAAAACCAT TGTCTACGAG GGTCCTGACG CGTGGGAGCT CTCTGGGATA CGTGTCACAC CTTCATCCTT CTGGAACGAG CTCTAGCCAA TGCCGGCAAT TGCTGGGCAAT GGTGTTCT GATCGTGGC CCACTGCTG GAGACAATCT AAATTATGAC AGAACAGCT CGAGACGTC CGACTGCTAG AGAACAGCT CACTGCTAG AGAACAGCT CACTGCTAG CACTGCTAG CACTGCTAG CACTGCTAG CACTGCTAG CACTGCTAG CACTGCTAGAC CTATGACAAC CTATGACAAC CTATGACAAC	51 AGCAAGATEG GGATACCAAC GTGCATCCGG CAGGCTTCCA GCGCTGGCCG GAGTGCGACT TGCCCGGCG CAGATGTACT AACTACGGGC GGAATAGTGG GTCGATATCT TTACCGTGTT GGTGAGACCG CACGTGTGCG CACGTGTCCG CACGTGTCCG CACGTGTCCG CACGTGTCCG CACGTGTCCG CACGTGTCCG CACGTGTCCT TCCAACACCA GACCTACTCA GCCTGCATTCT TCCAACACCA GCTGCCAAGG CTGATCACAC	60 120 180 240 300 360 420 480 540 660 720 960 900 960 1020 1140 1200 1260 1320
50 55 60	1 GTCATATTGA CTTTGAACTC CGGAAAACCC CTCAGTACTA ACCCCGTCGT AGAAAGCACT CTGGCCTACT CCTCAGGTAC CATCTCAGGAG GGGCGGCCTG ATGACAGAG GAGCACGAC TTAACAATCC CATCGGG AGAACAATGA AACCAGTGTG CCGGTGGGG CCGCTCCTGGT CCGGTGGGG AGAACAATGA AACCAGTGTG CCGGTGGGG CGCTCTCTCAGAT CAGCCATGAT CAGCCATGAT	11 ACATTCCAGA AGGGTCACCA CTATCCCGCA CCGTCCCC CTGCACGCAC CTGGAAGTTC CTGCATCACC GAATCGCTG GAAGTCCTGG CAGGGACATG ATOCACCAGC GTCACCACGC CATCACCCC CATCACCCC CATCACCCC CATCACCCC CATCACCCCC CATCACCCC CATCACCCCC CATCACCCC CATCACCCC CATCACCCC CATCACCCC CATCACCCC CATCACCC CATCACC CATCACCC CATCACCC CATCACCC CATCACC CATCACC CATCACC CATCACC CATCACCC CATCACC CATCAC CATCACC CATCACC CATCACC CATCACC CATCACC CATCACC CATCAC CATCAC CATCACC CATCAC CATCAC CATCACC	21 TACCTATCAT CAGCCACTG GTGCCCACTG GTGCCCAGT CCCAAATCCC TTGACCTGG ATGGCAGCA CCCTCTAACT CACCCTGTGT GGCTATAGA TTTATGARAC GATGCCTGTT TGGCTCA GAGTGGATCA GAGTGGATCA ACTCAAGCC CCAAAAGTCA ACGCAGTCA ACGCAGTCA ACGCAGTCA ACGCAGTCA ACGCAGTCA ACGAAAGGCA TTGAAAGCC CCAGGCATGA AGAAAGGGA AGAAGGGA AGATGCAACA TTCCTGCAGG	31 TACTCGATGC GACCTTACTA TGGTCCCCAC ACGCCCCAGA GCGCCCCAA GGTGTGATGG ACGGACCAA ATAATTTTTA TGAACACAA GCCAGAGCAG CCTGCACGT TGACAGCCGC CCGGGATTTTT TTTCTCATCC AGAAGCCTCT AGAAGCCTCA AGAAGCCTCA GCAGATATGT GCAGAGCTCA GCAGATATGT GCAGAGCTCA AGAACCTCAGA GCAGATATGT GCAGAACCTCAGA GCAGATATGT GGAACCTCAGA	41 TGTTGATAAC TGAAAACCAT TGTCTACGAG GGTCCTGACG AGTGTGCACC CTCTGGGATA CGTGTCACAC CTCTACCAC CTCATCCTT CTGGAACGAG CTCTAGCCAA AGTGGTTTACT GATCGTGGGC CCAGAACGTC CAGAACATCT AAATTATGAC AGAACAGCTC CTATGACAA CGCTGTGCAC CCACTGCTGC CAGAACATCT AAATTATGAC CACTTCTAAC AGAACAGCTC CTATGACAAC CTATGACAAC CTATGACAAC CTATTGCCAG	51 AGCAAGATEG GGATACCAAC GTGCATCCGG CAGGCTTCCA TCAAGACTA GCGCTGGCCG GAGTGCGACT TGCCCGGCG GAATAGTGG GTGATATCT TTACCGTGTT TTACGTGTT TTACGTGTTC TCAAGACCA GACTAGTCC GCAAAACCTC TCATGTTCT TCCAAGACCA GACCTAGTGA TGCTGCAAGG CTGCTACACC GCTGACACC GGTGACAGTG GGTGACAGTG GGTGACAGTG GGTGACAGTG GGTGACAGTG GGTGACAGTG	60 120 180 240 300 420 480 540 660 720 780 960 1020 1080 1140 1260 1320 1380
50 55 60	1 GTCATATTGA CTTTGAACTC CGGAAAACCC CTCAGTACTA ACACCGTACT CTGCCTACT CTGCCTACT CTGCCTACT CTGCCTACT CTGCCTACT ATGACAGCA ATAAAAACT TAGCCTGCG GAGCTCCAT TTAACAATCA ATGAGCCGG ACACCAGTGT CCGGGTGGGG TCCTCCAT GTGCACCATGAT CAGCCATGAT CAGCCATGAT CAGCCATGAT GAGGGCCTC	11 ACATTCCAGA AGGGTCACCA CTATCCCGCA CCGTCCCCC CTGCACGCAG GTGCATCACC CTGGAAGTTC CTGCATCACC GAATCACC GAATCACC GTACACACG GTACACACG GTACACACG GTACACACG GTACACACG GTACACACG CATCACCCC ATGCCATC ATGCCACAC ATGCCATC CTTGCCCAAC GGCCACCGAG TGAGACACAG TGCCAAAGCT TGCCAAAGCT	21 TACCTATCAT CCAGCTATTG CAGCCACTG GTGCCCAGT CCCAAATCCC TTGACCCTGG ATGGCAGCA CCCTCTAACT GACCCTGTGT GGCTATAAGA TTTATGAAAG TTTATGAAAG CAGGCAGTCA ACGGCATTG CAAAAAGTGA ATGAAGCTCC CCAGGCATGA AGGGAAGGA AGATGAACAACA TTCCTGCAGG AACAACAATA TACAGACCAG	31 TACTCGATGC GACCTTACTA TGGTCCCCAC ACGCCCCAGA CGTCCTCCT AGTGCTCCA ACGCCCAAA GCCAAGACGA ATAATTTTA TGAACACAAA GCCAGAGCAC GCCTGCACGC CCGCGGATTTT TTTCTCATCC AGACCTCAGA GCAGACGTC AGACCTCAGA GCAGACTCAGA CCAGACTCAGA CCAGACTCAGACA CCAGACTCAGACA CCAGACTCAGACA CCAGACTCAGACA CCAGACTCAGACA	41 TGTTGATAAC TGANAACCAT TGTCTACGAG GGTCCTGACG GGTCCTGACG CCTCGGGACCAT CCTCATCCAT CTGGAACGAG CTCTAGCCAA TGCCGGCAAT TGCCGGCAT CCACTGCGCC CACTGCGTC CACTGCGTC CACTGCGTC CACTGCGTC CACTGCGTC CACATCCTT AATTATGAC GACTATCAAC CTCTTGACAAC CTCTTGCAAC GACTTCTAAC CACTGCGTC CACTGCGTC CACTGCGTC CACTGCCTC CACTGCCT CACTGCCT CACTGCCT CACTGCCT CACTGCCT CACTGCC CACTGCCT CACTGCC CACTGCCT CACTGCCT CACTGCC CACTGCCT CACTGCC CACTGCCT CACTGCC CACTGCCT CACTGCC CA	51 AGCAAGATGG GGATACCAAC GTGCATCCGG CAGGCTTCCA GCGCTGGCCG GAGTGCGACT TGCCCGGCG GAGTGCGACT AACTACGGC GGAATAGTGA GTGGATATCT TACAGCTGTT GGTGAGAGCG CACATGTGCT TCATGTTCT TCCAAGACCA GACCTAGTGA TGCTGGATTT GCTGGATTT TCCAAGACCA GACCTAGTGA TGCTGGATTT GCTGGATTT GCTGGATTCT TCCAAGACCA GACCTAGTGA ACACTAGTGA TGCTGGATTT GCTGCCAAGG CTGATCACAC GGTGACAGG CTGATCACAC GGTGACAGG GTATTCACGG	60 120 180 240 300 360 420 480 540 660 720 960 900 960 1020 1140 1200 1260 1320
50556065	1 GTCATATTGA CTTTGAACTC CGGAAAACCC CTCAGTACTA ACCCCGTCGT AGAAAGCACT CTGGCCTACT CTGGCCTACT CGGAGGACGA GGGCGGCCTG ATGACAGCGG ATAAAAAACT ATGCCTGCGG GAGGCTCCAG AGAACAATGA AACCAGTGTC CCGGGTGGG TGCTTCTCAT CAGCCATGAT CAGCCATGAT CAGCCCTTCAT CAGCCATGAT CAGCCCTTCAT CAGCCATGAT CAGCCCTTCAT CAGCCATGAT CATGGATTTA	11 ACATTCCAGA AGGGTCACCA CTATCCCGCA CCGTCCCCC CTGCACGCAG GTGCATCACC CTGCATCACC CTGCATCACC CTGCATCACC CTGCATCACC CTGCATCACC CAGGGACATG ATCCACCAGT GGTCAACTTG GGCCTGGCCC CATCACCCC CATCACCCC CATCACCCC CATGCCATGG ATACCACAGT CGCCAACGT TCGCCAAC TGGCCACGAG TGAGACACAG TGAGACACAG TGAGACACAG TGCCAAAGCT TCCGACAAAGCT TCCGACAAATT CTGCCAAAGCT TCCGACAAATT	21 TACCTATCAT CAGCTATTG CAGCCACTG GTGCCCAGT CCCAAATCCC TTGACCTGG ATGGCAGCA CCCTCTAACT GTTCGCCTCT CACCTGTGT GGCTATAGA TTTATGAAAC GATGCCTGTT AACTCAAGCC TGGCAGGTCA ACGCAGTTG CAAAAAGTGA ATGAAAGTGA ATGAAAGGAA AGATGAACA TTCCTGCAGG AACAACAATA TACAGACCAG AAGGCAAAC AAGGCAAACCA TACAGACCAG AAGGCAAACC	31 TACTCGATGC GACCTTACTA TGGTCCCCAC ACGCCCCAGA GCGCCCCAA ACGCCCCAA ACGCCCCAA ACGCCCCAA ACGCCCCAA GCTACTCCT AGTGCTCCA AGTGCTCCAA GCCAAGACGA ATAATTTTA TGAACACAGA GCCAGAGCAG GCCTGCACGT TGTCTCATCC AGAAGCCTCT TGTTCTCATCC AGAAGCTCT TGCTGCAGC GCAGATATGT GGAACGTCGA GCAGATATGT GGACGTCGA GCAGTTACCA GCAGTTACCA GCAGTTACCA GCAGTTACCA GCAGTTACCA GCAGTTACCA GCAGTTACCAC GCAGTTACCAC GCCTAATCCAC	41 TGTTGATAAC TGAAAACCAT TGTCTACGAG GGTCCTGACG GGTCCTGACG CCTCGGGATA CCTCTCATCCT CTGGAACGAG CTCTAGCCAA TGCCGGCAAT TGCCGGCAAT TGCTGGGACGT CCAGCACGCT CAACTCCTC CAACTCCTC AAATTATGAC AGAACAATC AGTGCTGAAC CTATGACAAC CTATGACAAC TTCTGCCAG GATAGGCGAA TTCTTGCCAG GATAGGCGAA TCTTGCCAG GATAGGCGAT AAATGGCGAT CAATGGTCTCG CAACGCTCTCC	51 AGCAAGATEG GGATACCAAC GTGCATCCGG CAGGCTTCCA TCAAGACTA GCGCTGGCCG GAGTGCGACT TGCCCGGCG CAGATGTACT AACTACGGGC GGAATAGTGG GTGGATATCT TTACCGTGTT GGTGAGACCG CACGTGTCCG CACGTGTCCG CACGTGTCCG CACGTGTCCG CACGTGTCCG CACGTGTCCG CACGTGCACAC GCCTACGACCA GCCTACGACCA GCCTACGACCA GCTGCAAGG CTGATCACAC GCTGACAGG CTGATCACCG GTATCACGG TCCTTCACGT	60 120 180 240 360 420 480 540 660 720 960 960 1020 1140 1260 1320 1380 1440 1560
50 55 60	1 GTCATATTGA CTTTGAACTC CGGAAAACCC CTCAGTACTA ACCCCGTCGT AGAAAGCACT CTGGCCTACT CCTCAGGTAC GGGAGGACGA CATCTCAGAG GGCCGCCTG ATGACAGAG CGCTCCCGGG GAGGCTCCAT TTAACAATCC ATGAGACGGG AGAACAATGA AACCAGTGTG CCGGGTGGGG CGCTCTCTCAT CAGCCATGAT GAGGGCCTCT GTTCTCGCT GTTCTTCGCT CTTTTCTGCTT CCTTTTTCACA	11 ACATTCCAGA AGGGTCACCA CTATCCCGCA CCGTCCCC CTGCACGCAC CTGGAAGTTC CTGCATCACC CAGGACTCACC CAGGGACATG ATCCACCAG CTACCACCAGT CATCACCAGT CATCACCAGT CATCACCAGT CGCCTGCCC CATCACCCC CATCACCCC CATCACCCCC CATCACCCCC CATCACCCCC CTGCCCAACT CGCCACCGAG CTGTGCCGAC CTGTGCCGAC CTGTGCCGCG CTGCCAAACT TCGCCAAACT TCGACAAATG CAAAACAATG CAAAACAATG	21 TACCTATCAT CAGCCACTG CAGCCACTG GTGCCCCAGT CCCAAATCCC TTGACCCTGG ATGGCAGCA CCCTCTAACT CACCCTGTGT GGCTATAGA TTATGAAAC GATGCCTGTT AACTCAAGCC TGGCAGGTCA AAGACAACA AGAGAAAAGGA AGAGAAACACA TTCCTGCAGG AAGACAACAC AAGGCAAACG GGGCTGTTT TACAAACCA CAAAAACTGA ATCCTGCAGG AACAACAACA TTCCTGCAGG AACAACACA CGGCCAACCA GGGCAAACCA CGGCCAACACA CGGCCAACACA CAGGCCAACAC CGGCCAACAC CGGCCAACAC CGGCCAACAC CGGCCAACAC CGGCCAACAC CGGCCAACC CCCAACC CCCAACCAC CCCAACACC CGGCCAACC CGGCCAACC CGGCCAACC CGGCCAACC CCCAACC CCCCAA	31 TACTCGATGC GACCTTACTA TGGTCCCCAC ACGCCCCAGA GCGCCCCAA GGGCCCAAA ACGCCCCAA GCGACCAAA ACGGACCAA ACGGACCAA ACGGACCAA ACGGACCAC TTCAAAAGC GCCAGAGCAG GCCAGAGCAG GCCTGCACGT TGACAGCCGC AGAAGCTCA AGAAGCCTCA AGAAGCCTCA AGAAGCCTCA AGAAGCTCAA GCAGATATGT GGAACGTCAGA GCAGATATGT GGAACGTCAGA GCAGATATGT GGAACGTCAG GCAGATATCCAC GGATGTACCG GCTAATCCAC TGCTTCCCCG	41 TGTTGATAAC TGAAAACCAT TGTCTACGAG GGTCCTGACG GGTCTGACG CGTGGGACC CTCTGGGATA CGTGTCACAC CTTCATCCTT CTGGAACGAG CTCTAGCCAA TGCCGGCAAT AGTGGTTTCT GAACACGCTC GAGACAATCT AAATTATGAC AGAACAGCTC CTATGACAAC CTATGACAAC CTATGACAAC TTCTTGCCAG GATAGGGGAT TCTTGCCAG GATAGGGGAT GAATGTGATC ATGGTCTTCC TGCATGATT TGCCAG TGCATGATT TGCCAG TGCATGATT TGCCATGATT TGCATGATT TGCCATGATT TGCATGATT TGC	51 AGCAAGATEG GGATACCAAC GTGCATCCGG CAGGCTTCCA TCAAGACTA GCGCTGGCCG GAGTGCGACT TGCCCGGCG GAATAGTACT TTACGCTGTT TTACGCTGTT TTACGTGTT TCAAGACCA GACTAGTCC TCATGTTCT TCCAAGACCA GACTAGTCA GTGAAGTC GCTGATCACAC GGTGACAGT ACTACTTCACGT TCCTTGACGT ACTCTTAGAG	60 120 180 240 360 420 480 540 660 720 840 900 1020 1080 1140 1260 1320 1320 1340 1500
50556065	1 GTCATATTGA CTTTGAACTC CGGAAAACCC CTCAGTACTA ACCCCGTCGT AGAAAGCACT CTGGCCTACT CTGGCCTACT CTGGCCTACT CTGGCCTACT CTGGCCTACT CTGGCCTACT CTGGCCTGG GGGGGGCCTG ATGACAGCG GAGGCTCCAG ATGACAGTAC ATGCGGTGGG TGCTTCTCAT CAGCCATGAT CAGCCATGAT CTTTTTACAA ACCAGTGT CTTCTGGCTG ACTGGATTTA CATTTTTACAA ATGATTCAGA TGCCATACTG	11 ACATTCCAGA AGGGTCACCA CTATCCCGCA CCGTCCCCC CTGCACGCAG GTGCATCACC CTGGAAGTTC CTGCATCACC CAATCGCTG GAAGTCCTGG CAGGGACATG GTCCACCAGT GGTCAACTTG GGCTACACTTG GGCATGACTTG CATTGCCACC CATCCCCC ATACCACCCC ATACCACCCC ATACCACCCC ATACCACCCC ATACCACCCC TCTGCCAAC TCTGCCAAC TCTGCCAAC TGGACACTGG TCCCAACGTG TCCCAACGCTCG TCCCAAAGCT TCCCAAAGCTT TCCAACACTC GAAAACAATG GAAAACAATG GACACATCC GGTCACTTCA TCGACAAATC TTCCACCACC GGTCACTTCA TCGACAAATC TTCCACCACCTCC TTCCACACCTCC TTCCACCCTCC TTCCACCTCC TTCCACCCTCC TTCCACCTCC TTCCACCTCC TTCCACCCTCC TTCCACCCTCC TTCCACCTCC TTCCACCTC TTCC	21 TACCTATCAT CAGCTATTG CAGCTATTG CAGCCACTG GTGCCCAGT CCCAAATCCC TTGACCTG ATGGCAGCA ATGGCAGCA ATGGCAGCA GATCCCTGTAACT TATATGAAAC GATGCATTG AACTCAAGCC TGGCAGTCA AAGTGATCG AAGAAAGTGA ATGAAAGTGA ATGAAACACAATA TACAGACCAG AAGGCAACA AAGGCAAACG GGCTGGTTT TACTAAACA TTCTTGCAGG AAGAAAACA TTCTTATTAA AAGGCCACTC TTTTTTTTTT	31 TACTCGATGC GACCTTACTA TGGTCCCCAC ACGCCCCAGA CGTCCTCCT AGTGCTCCT AGTGCTCCAA ACGCACAAA CCCAAGACGA ATAATTTTA TGAACACAAG CCCAGAGCAG GCCTGCACGT TGACAGCCGT TGACAGCCGT AGACCTCAGA GCAGATATGT TTCTCATCC AGAACTCAGA GCAGATATGT GGAACGTCGA GCAGATTGT GGAGCTCCAGC TCTGCAGC TCTGCAGC TCTGCAGC TCTGCAGC TCTGCAGC TCTGCAGC TCTGCAGC TCTGCAGC TCTGCAGC TCTTCCCCC TCTTCCCCC TCTGCCCCACC CTGCCCCACC CCCCCCCC	41 TGTTGATAAC TGAAAACCAT TGTCTACGAG GGTCCTGACG CGTGGGAGCT CTCTGGGATA CCTCATCCTC CTGGAACGAG CTCTAGCCAA TGCCGGCAAT TGCCGGCAAT TGCCGGCAAT CCATCCTC CAATCCTC CAATCCTC CAATCCTC CAATCCTC CAATCCTC AATTATGAC GACATCTTCAAC CTATGACAAC CTATGACAAC CTATGACAAC CTATGACAAC TCTTTGCCAG GAAACGTC AATGACAAC TCTTGCCAG GAATGTGATC TGCATGATTT TGCTCGGCTT TGCCTCCCT TGCATCCTCCT TGCATCCTCCT TGCATCATTT TGCTCCCTT TGCATCACT TGCATCACTT TGCATCACT TGCATCA	51 AGCAAGATGG GGATACCAAC GTGCATCCGG CAGGCTTCCA GCGCTGCCCG GAGTGCGACT TGCCCCGGCG CAGATGTACT AACTACGGC CAGATGTACT TTACCGTGTT CGTGAGACCG CACATGTTCT TCCAAGACCA GACCTACTCT TCCAAGACCA GACCTACTGGTGACACG CTGATATCT TCCAAGACCA GACCTACTGGATTT GCTGCCAAGG CTGATCACAC GGTGACAGT CTGATCACAC GGTGACAGT CTGATCACAC GGTGACAGT CTGATCACAC GGTGACAGT CTGATCACAC GGTGACAGT TCCTTGACGT TCCTTGACGT ACTCTTTAGAG TGGCACTTCT AACCCCTTGT	60 120 180 240 360 420 480 540 660 720 900 900 1020 1080 1140 1260 1320 1320 1560 1560 1680 1740
50556065	1 GTCATATTGA CTTTGAACTC CGGAAAACCC CTCAGTACTA ACCCCGTCGT AGAAAGCACT CTGGCCTACT CTGGCCTACT CGGAGGACGA GAGGACGA ATAAAAACT TTAACAATCA ATGACTGCC CGGTGGG AGACAATGA AACCAGTGTC CCGGGTGGGG TGCTTCTCAT CAGCCATGAT CAGCCATGAT CATCTGATTA CATTTACAA ATGATTCAA ATGATTCAA ATGATTCAA ATGATTCAA ATGATTCAA TCGCATACT CCCCAAAGGG CCCCATACT CCCCCAAGGGG CCCCATACT CCCCCAAGGGG CCCCATACT CCCCCAAGGGG	11 ACATTCCAGA AGGGTCACCA CTATCCCGCA CCGTCCCCC CTGCACGCAG GTGCATCACC CTGCATCACC CTGAAGTTC CTGCATCACC GAATCGTGT GAGTCCTGG CAGGGACATG ATCCACCAGT GGCACTAGC CATCACCCC CATCACCCC CATCACCCC CATCACCCC CATCACCCC CATCACCCC CATCACCCC CATGCCAACT GGCCACCGAG TGAGACACAG CTGTGCCAAC CTGTCCCAAC CTGTCCAAACT TCGACAAACT TCGACAAACT TCGACAAACT GGTCACTTCA GGTCACTTCA GGTCACTTCA CATGCCAAACC TTCGACAAACC TCGACAAACC TTCGACAAACC TTCGACAACC TTCGACACC TTCGACCACC TTCACTCC TTCACCACC TTCACTCC TTCACCCC TTCACCCC TTCACCCC TTCACCCC TTCACCCC TTCACCCC TTCACCCCC TTCACCCCC TTCACCCC TTCACCCCC TTCACCCC TTCACCCC TTCACCCCC TTCACCCC TTCACCC TTCACCCC TTCACCC TTCACC TTCACCC TTCACC TTCACCC TTCACCC TTCACCC TTCACCC TTCACCC TTCACCC TTCACCC TTCACCC	21 TACCTATCAT CAGCCACTG CAGCCACTG GTGCCCAGT CCCAAATCCC TTGACCCTGG ATGGCACAC CCCTCTAACT GTTCGCCTCT GTTCGCCTCT GTTCGCCTCT GTTCACCT GTTCACCT GAGCAGACA CATCCAAGCC TGGCAGGTCA ACTCAAGCC CAAAAAGTCA ATGAAAGTCA ATGAAAGTCA TTCCTGCAGG AACACAATA TACAGACCA GAGCAACAC TACCTGCAGG AACACACATT TACTACACC GGCTGGTTT TATTTATTAA AAGGCAAACC GGCTGGTTT TTTTTTTTTT	31 TACTCGATGC GACCTTACTA TGGTCCCCAC ACGCCCCAGA GCGCCCCAGA GGGCCTCCAA GCCAGAGACGA ATAATTTTTA TGAACACAGA GCCAGAGCCAC GCCGGGATTTTTTCTATCC AGAGCCTC AGACCTCCA AGACCTCC AGACCTCCAC ACAGTCACC CCTGCACCC CCAGTCACC CCAGTCACC CCTCCCACC CCAGTCACC CCTCCCCCC CCACTCCCCC CCACTCCCCC CCACTCCCCC CCACTCCCCC CCACTCCCCC CCACTCCCCC CCACTCCCCC CCACTCCCCC CCACTCCCCC CCACTCCCCCC CCACTCCCCCC CCACTCCCCCC CCACTCCCCCC CCACTCCCCCC CCACTCCCCCC CCACTCCCCC CCACTCCCCC CCACTCCCCCC CCACTCCCCCC CCACTCCCCCC CCACTCCCCC CCACTCCCCCC CCACTCCCCC CCACTCCCCCC CCACTCCCCC CCACTCCCCCC CCACTCCCCCC CCACTCCCCCCC CCACTCCCCCC CCACTCCCCCC CCACTCCCCCC CCACTCCCCCC CCACTCCCCCC CCACTCCCCCC CCACTCCCCCC CCACTCCCCCCC CCACTCCCCCC CCACTCCCCCC CCACTCCCCCC CCACTCCCCCC CCACTCCCCC CCACTCCCCC CCACTCCCCC CCACTCCCCC CCACTCCCCC CCACTCCCCC CCACTCCCCC CCACTCCCCC CCACTCCCCC CCACTCCCCCC CCACTCCCCC CCACTCCCC CCACTCCCC CCACTCCCC CCACTCCC CCACTCCCC CCACTCCC CCACTCCC CCACTCCC CCACTCCC CCACTCCC CCACTCCC CCACTCCC CCACTCCC CCACTCCC CCACTCC CCACTC CCAC	41 TGTTGATAAC TGAAAACCAT TGTCTACGAG GGTCCTGACG GGTCCTGACG CCTGGGAGCT CTCTGGGATA CGTGTCACAC CTTCATCCTT CTGGAACGAG CTCTAGCCAA TGCCGGCAAT TGCCGGAACGTC CCACTGCGTG GAGACAATCT AAATTATGAC AGAACAGCT CTATGCAAC CTATGACAAC TCTTGCCAG GATAGGGAT TCTTGCCAG GATAGGGAT TCTTGCCAG TGATGTTTCT TGATCTCCCT TCCAATTCTG	51 AGCAAGATEG GGATACCAAC GTGCATCCGG CAGGCTTCCA CAGCCTGGCCG GAGTGCGACT TGCCCGGCG CAGATGTACT AACTACGGGC GGAATAGTGG GTCGATATCT TTACCGTGTT GGTGAGACCG CACGTGTCCG CACGTGTCCG CACGTGTCCG CACGTGTCCG CACGTGTCCG CACGTGTCCACGC CTCATCTTCT TCCAACGCCATGC CACGTGTCACG GTGACACTC GCTGCATTCACGC ACACCTCGGT ACTCTTCACGT ACCCCTTCT AACGCACGG CACCCCTTCT AACGCACGG	60 120 180 240 360 420 480 540 660 720 960 900 910 1080 1140 1260 1320 1380 1440 1560 1560 1620 1680 1680 1680 1680
5055606570	1 GTCATATTGA CTTTGAACTC CGGAAAACCC CTCAGTACTA ACCCCGTCGT AGAAGACACT CCTCAGGTAC GGGAGGACGA CATCTCAGGG ACATCTCAGAG GGCCGCCTG ATGACAGAG CGCTCCCGGG GAGGCTCCAT TTAACAATCC ATGAGAGCGG AGAACAATGA AACCAGTGTG CCGGGTGGGG TGCTTCTCAT CAGCCATGAT CAGCCATGAT ACTTTTACAA ATGATTCAGA ATGCATTTA	11 ACATTCCAGA AGGGTCACCA CTATCCCGCA CCGTCCCC CTGCACGCAC CTGCATCACC CTGCATCACC CTGCATCACC CTGCATCACC CTGCATCACC CTGCATCACC CTGCATCACC CAGGGACATG ATOCACCAGT GGTCAACTTG GGCCTGGCCC CATCACCCC CATCACCCCC CATCGCCTAC CGCCACCGAC TGCCCACC GGCCACCGAC TGGCCACCGAC TGGCCACACT TGCCCAAACT TGCACAATT CAACACTT CAACACTT CAACACTT CAACACT TGCCCAAACC TTCGACCAAAT GGTCACTTCA CAACACT TCGACCAAAT TGCACGAC TGCCAACCT TGCACACT TGCACGCC TGCCCACTTC TGATGCCGG GCCCCCATTC	21 TACCTATCAT CAGCCACTG CAGCCACTG GTGCCCCACTG GTGCCCCACTG TTGACCCTGG ATGGCCACCA CCCTCTAACT CACCCTGTGT GGCTATAGA TTTATGABAC GATGCCTGTT AACTCAAGCC TGGCAGGTCA AAGACAACA AGAGAAAGGA AGAGAAACACA TTCCTGCAGG AAGACAACA TTCTTGCAGC AAGGCAACCA CGGCTGTTT TATTATAAAAC CCCGGCTGTTT TACTGCAGG AACACACA CCCGGCTGTTT TTTTTATTAA AGTGCATCCC CTGGTTGTGG AGATCTCCC CTGGTTGTGG AGATCTCCT CTGGTTGTGG AGATCTTCCT CCCGGTTGTTGG AGATCTTCCT CCAGGTTGTTGG AGATCTTCCT	31 TACTCGATGC GACCTTACTA TGGTCCCCAC ACGCCCCAGA GCGCCCCAA GGGACCAAA ACGGACCAAA ACGGACCAAA ACGGACCAAA ACGGACCAAA ACGACACAC ATAATTTTA TGAACACAGG GCCAGAGCAG GCCTGCACGT TGACAGCCGC AGACCTCAGA GCAGATATT TTTCTCATCC AGAAGCCTCT TGCTGCAGC GCAGATATGT GGAACGTCGA GCAGATATGT CAGAGCTCCA ACAGTGACCC TCTCCCCG ACAGTGACCT CTGCCCAGC CCAGTGACCT CTGCCCAGC CCACTGCCCG CCACTGCCCG CCACTGCCCG CCACTGCCCG CCACTGCCCG CCCACTGCCCG CCCTAGTCCCT CCCCACCC CCCTGCCCGCC CCCTGCCCGCC CCCTGCCCGCC CCCTGCCCGCC CCCTGCCCACCC CCCTGCCCGCC CCCTGCCCACCC CCCTGCCCACC CCCTGCCCACC CCCTGCCCGCC CCCTCCCCACC CCCTCCCCACC CCCTCCCCACC CCCTCCCCACC CCCTCCCCACC CCCTCCCCACC CCCCCCCC	41 TGTTGATAAC TGAAAACCAT TGTCTACGAG GGTCCTGACG GGTCCTGACG CGTGGGACCT CTCTGGGATA CGTGTCACAC CTTCATCCTT CTGGAACGAG CTCTAGCCAA TGCCGGCAT AGTGGTTCAC CCACTGCTG GAACAACTC AAATTATGAC AGAACAGCT CTATGCAAC CTATGACAAC TTCTTGCCAG GATAGGGAT TCTTGCCAG GATAGGGAT TCTTGCCAG TGAATCTTAC TGCATGATT TGCTTCCCT TGCATGATT TGCTCTCCCT TCCATGGGG TTCCAGGGG TTCCAGGGG TTCCAGGGG TTCCAGGGGC	51 AGCAAGATEG GGATACCAAC GTGCATCCGG CAGGCTTCCA TCAAGACTA GCGCTGGCCG GAGTGCGACT TGCCCGGCG GAATAGTACT TTACCGTGTT TTACGTGTT TTACGTGTT TCAAGACCA GACTAGTCA ACCCTTGT ACTCTTAGAG TGCTTTAGAG TGCCTTTAGAG TGCCCTTTC AACCCCTTCT AACCCCTTCT AACCCCTTCT AACCCCTTCT AACCCCTTCT AACGCACGG CAATTTTGGA	60 120 180 240 300 420 480 540 660 720 780 960 1020 1140 1260 1320 1380 1440 1560 1680 1740 1860
50556065	1 GTCATATTGA CTTTGAACTC CGGAAAACCC CTCAGTACTA ACCCCGTCGT AGAAAGCACT CTGGCCTACT CTGGCCTACT CTGGCCTACT CTGGCCTACT GGGAGGACGA GGGCGGCCTG ATGACAGCGG ATGACAACA TAGCCTGCGG GAGCTCCAT ATTGACAGCG CACTCCT ATGAGCGG TCACTGGT CAGGCTCCAT ACCAGTGT CAGGCCTCAT CAGCCATGAT CAGCCATGAT CAGCCATGAT CAGTTTTACAA ATGATTCAGA ATGATTCAGA TGCCATACTG CCGCAAGGGG CTTGGAGGC TTGAGGGCTCCT CCGCAAGGGG CTTGGAGGCT TGAGGCATGT CCGCAAGGGG GTTGGAGGAGGAGGAGGAGGAGGAGGAGGAGGAGGAGGAG	11 ACATTCCAGA AGGGTCACCA CTATCCCGCA CCCGTCCCCC CTGCACGCAG GTGCATCACC CTGGAAGTTC CTGCATCACC GAATCGCTG GAAGTCCTGG CAGGGACATG GTCAACTG GTCAACTG GTCAACTG GTCAACTG GTCAACTG CATGCCCC CATCACCCC CATCCCACC GGTCAACTG TCGCCAAC TGGCAAC TGGCAAC TGGCAAC TGGCAAC TGGCAAC TGCAAAGCT TCGACAAGCT TCGACACTGC GTCACTTCA TCAGGCTGC TGATGCCGG GCCCCCATTG GCTGCCCGT ACAGCCAGGT ACAGCCAGGT ACAGCCAGGT	21 TACCTATCAT CAGCTATTG CAGCTATTG CAGCCACTG GTGCCCAGT CCCAAATCCC TTGACCTGG ATGGCAGCA ATGGCAGCA ATGGCAGCA GATCCCTGTTG GACCTGTTAACA TATATGAAAC CAGCAGTCA ACGCAGTCA ACGCAGTCA ACGCAGTCA ACGCAGTCA ACGCAGTCA ACGCAGTCA ACGCAGTCA ACGCAGTCA TCCTGCAGG AGAAAAGGA AGATGAACA TTCTTGCAGG GACAACA TTCTTTATTAA AGTGGCTCC CTGGTTGTG AGATCTCCT CTCAGCTCC CTGGTTGTG AGATCTCCC CTGGTTGTG AGATCTCCT CTCAGCTCCC CTGGTTCTCG GGCACCTCCA	31 TACTCGATGC GACCTTACTA TGGTCCCCAC ACGCCCCAGA CGACCTTCCT AGTGCTCCAA ACGCCCAAA ACGCCCAAA GCCAAGACGA ATAATTTTA TGAACACAAG CCCAGAGCAG GCCTGCACGT TGACAGCCGC AGACCTCAGA GCAGATATT TTTCTCATCC AGAACTCAGA GCAGATATGT GGAAGCTCA GCAGTGAACC CCAGGTGACC TCTCCCCG GCTTCCCCC GCAGTGACC CCACTGCACC CCACTGCACC CCACTGCACC CCACTGCACC CCACTGCACC CCACTGCACC CCACTGCACC CCACTGCACC CCACTGCACC CCACTGCCC CCACTGCCC CCACTGCCC CCACTGCCC CCACTGCCC CCACTGCCC CCACTGCCC CCACTGCCCC CCACTGCCCC CCACTGCCCC CCACTGCCCC CCACTGCCCC CCACTGCCCC CCACTCCCC CCACTCCCCC CCACTCCCCC CCACTCCCC CCACTCCCCC CCACTCCCCC CCACTCCCCC CCACTCCCC CCACTCCCCC CCACTCCCCC CCACTCCCCC CCACTCCCCC CCACTCCCCC CCACTCCCC CCACTCCCCC CCACTCCCCC CCACTCCCCC CCACTCCCCC CCACTCCCCC CCACTCCCCC CCACTCCCCC CCACTCCCCC CCACTCCCC CCACTCCCCC CCACTCCCCC CCACTCCCCC CCACTCCCCC CCACTCCCCC CCACTCCCCC CCACTCCCCC CCACTCCCC CCACTCCC CCACTCC CCACTC CCAC	41 TGTTGATAAC TGAAAACCAT TGTCTACGAG GGTCCTGACG GGTCCTGACG CGTGGGACGT CTCTGGGATA CGTGTCACAC CTTCATCCTT CTGGAACGAG CTCTAGCCAA TGCCGGCAAT TGCCGGCAAT TGCCTGGGACGTC CAATCGTG CAATCTTCAC CAATCGTG AATTATGAC GACTTCTAGCAA CTCTTGCCAG GATAGGGAT TGCTGGCAG TTCTGCCT TGCATGTTCT TGCTCTCCT TCCATGCTT TGCTCCCT TCCAGTGCTT TGCTCCCT TCCAGGGCC TCCAGTGCAAC TTCTGCCTT TGCTCTCCCT TCCAGTGCAT TCCAGGGGCC AGATGAAAA TCTGGGGCCA	51 AGCAAGATTG GGATACCAAC GTGCATCCG GGATACCAC GTGCATCCG CAGGCTTCCA GCGCTGCCC GAGTGCGACT TGCCCGGCG CAGATGTACT TACCGGC GGATAGTT TTACGGTTT GGTGAGACC CACGTGTCC GAAAACCTC TCATGTTC TCCAAGACCA GACCTAGTGA TCCTAGTATT GCTGCCAAGG CTGATCACAC GGTGACAGT CTGATCACAC GGTGACAGT TCCTCTAAGG TCCTTTAAGG TCGCTTCT AACCCCTTGT AAGCAGAGAC CAATTTTGGA GGAGACAT CTTGGTAGTG CATTTGGTAGTG	60 120 180 240 360 420 480 540 660 720 960 900 910 1080 1140 1260 1320 1380 1440 1560 1560 1620 1680 1680 1680 1680
5055606570	1 GTCATATTGA CTTTGAACTC CGGAAAACCC CTCAGTACTA ACCCCGTCGT AGAAAGCACT CTGGCCTACT CTGGCCTACT CTGGCCTACT CTGGCCTACT AGAAAGCACA GGGAGGACGA ATAAAAAACT TTAACAATCC ATGGACTCAGG AGACCAGGT CCGGGGGGCTCAT TTAACAATCA ATGACAGTGTC CTGTCTCACT CAGCCATGAT CAGCCATGAT CAGCCATCAT ACTGGATTTA CGTTTTACAA ATGATTCAGA ATGATTCAGA ATGATTCAGA TGCATTACAT CCCCAAGGGG CTTGGAGGCT CTGGAGGCT TTAGACATCG CTGGAGGGCT TCAGCCATGAT CCCCAAGGGG TTCGCAAGGGG TTCGCCAGCCT TCAGCCATGAT TCAGCATGAT CCCCAAGGGG TTCAGCATGGA TCCCCAGCCT	11 ACATTCCAGA AGGGTCACCA CTATCCCGCA CCGTCCCCC CTGCACGCAG GTGCATCACC CTGCATCACC CTGCATCACC CTGCATCACC CTGCATCACC CTGCATCACC CTGCATCACC CAGGGACATG ATCCACCAGT GGTCACTAGC CTACCACAGT GGCCACCGAG TCACCACCAC TCGCCAACTTC GGCCACCGAG TGAGACACAG TGAGACACAG TGCCAAACTT CGCCAAACTT CGCCAAACT TCGACAAATG GGTCACTTCG GGTCACTTCG GGTCACTTCG CTGCCCAACGC TGCCAAACGC TGCCAAAGCT TCGACAAATG GAAACAATG GGTCACTTCG GCTCACTTC GCTGCCCATTC CCACCGGT ACACCCAGGT ACACCCAGGT ACACCCAGGT ACACCCAGGT ACACCCAGGT ACACCCAGGT ACACCCAGGT ACACCCAGGT ACACCCAGGT ACTCACAAAG	21 TACCTATCAT CAGCCACTG CAGCCACTG GTGCCCAGT CCCAAATCCC TTGACCCTGG ATGGCAGCA ACCCTGTGT GGCTATAACA TTTATGARAC GATGCCTGT AACTCAAGCC TGGCAGGTCA ACGCAGTTG CAAAAAGTCA ATGAAAGTCA ATGAAGCC TGGCAGGATTG CAAAAAGTCA ATGAAGCACA ATCCTGCAGG AACACAATA TTCTGCAGG AACACACAC TTCTTTATTTATTATTATTATTATTATTATTATTATATATATA	31 TACTCGATGC GACCTTACTA TGGTCCCCAC ACGCCCCAGA GCGCCCCAGA GGACCTTCCT AGTGCTCCAA GGTGGACCAAA GCCAAGACGA ATAATTTTTA TGAACACAGA GCCAGAGCCAG GCCAGAGCCAG GCCAGAGCCG GCCAGCCCAG GCAGCTGCCC TGATGCTGCAGC GCAGTGACCC GCAGTGACCC TGCAGCC GCAGTGACCC TGCAGCC GCAGTGACCC TGCAGCC GCAGTGACCC GCACTGCCG GCACTGCCG GCACTGCCG GCACTGCCC TGATGCCC TGATGGCTC TGATGGCTC TGCCCAGCC GCACTGCCC GCACTGCCC TGCATGCCC TGCCCAGCC TGATGCCTC TGATGGCTC TGATGGCTTC	41 TGTTGATAAC TGAAAACCAT TGTCTACGAG GGTCCTGACG GGTCCTGACG CCTGGGAGCT CTCTGGGATA CCTTCATCCTT CTGGAACGAG CTCTAGCCAA TGCCGGCAAT TGCCGGCAAT TGCTGGGACGTC GAACGTC CAATGCTAA CAATTATACAC GAATTATACAC GAATTATACAC TCTTGCCAG GATAGAGAAC TCTTGCCAG GATAGGGAT TGCTTGCCAG TCTTGCCAG TCTTGCCAG TCTTGCCAG TCTTGCCAG TCTTGCCTT TCAATTTTCG TCCATGCTT TCCATGCTT TCCATGCTT TCCATGCTT TCCATGCCT TCCATGCCAT TCCAGGGC AGATGAAAAC TCTTGCCAT TCCAGGGCC TCCAGGGCC TTAGAGCCTT	51 AGCAAGATEG GGATACCAAC GTGCATCCGG CAGGCTTCCA GCGCTGGCCG GAGTGCGACT TGCCCGGCG CAGATGTACT AACTACGGC GGAATACTA GCGCTGCCG GGAATACTG GTGGATATCT TTACGGTGT GGTGAGACC CACGTGTCC GAAAAACCTC TCATGTTCT TCCAAGACCA GACCTAGTCA GCCTGCATG GCTGCATG CTGCTACAC GCTGCATG CCTGCTACAC GCTGCATG ACCCTGTTCACGT ACCCTTTACAG TCCTTTACAG TCTTACAG TCTTCTTACAG TCTTCTTACAG TCTTCCTTACAG TCTTCCTTACAG TCTTCCTTACAG TCTTCCTTACAG TCTTCCTTACAG TCTTCCTTCTACAG TCTTCTTCTACAG TCTTCTTCTACAG TCTTCTTCTACAG TCTTCTTCTACAG TCTTCCTTCTACAG TCTTCTTCTACAG TCTTCTTCTACAG TCTTCTTCTTCTACAG TCTTCTTCTTCTACAG TCTTCTTCTTCTTCTTCTTCTTCTTCTTCTTCTTCTTCT	60 120 180 240 360 420 480 540 660 720 960 900 910 1080 1140 1260 1320 1380 1440 1560 1560 1620 1680 1780 1860 1980 1980 1980
5055606570	1 GTCATATTGA CTTTGAACTC CGGAAAACCC CTCAGTACTA AGAAGCACT CTGCCTAGTACT AGAAAGCACT CTGCCTAGTAC CGGAGGACGA CATCTCAGAG GGGGGGCCTG ATAAAAACT TAGCCTGCG ATAAAAACT TAGCCTGCGA AGAACAATC ATGGAGCCG AGAACAATC ATGACTGCT CAGGGTGGG TGCTTCTCAT CAGCCATGAT CAGCCATGAT CAGCCATGAT CAGCCATGAT CAGCCATGAT CAGCCATGAT CAGCCATGAT CAGCCATGAT CAGCCATGAT CAGCCATACTG CCCAAGGG CTTGGAGTTA ATGATTCAGA TGCCATACTG CCCCAAGGG CTTGGAGGC TGAGCATGA TCAGCCC TGAGCATGA TCAGCCC TGAGCATGA TCAGCC TGATGTGGC TCAGTGGCC TGAGGAGGC TCCCCAGCC TGATGTGGCC TGATGTGGCC	11 ACATTCCAGA AGGGTCACCA CTATCCGGCA CTATCCCGCA CTGCACCCC CTGGAAGTTC CTGCATCACC CTGGATCACC CTGGATCACC CTGGATCACC CTGCACCAGG GAACGCTGC CAGGACATTG GATCACCACAGT GGCACACAGT CATCACCCC ATGCCTTGCCCACACCCC ATGCCACACCCC ATGCCACACCC GGCCACCGAG CTGTCCCGC GGCCACCGAG CTGTCCCGC TCGACAACTTC TCGACAACTTC GGCACAATTC GGTCACTTCACCCC TGCACACTTCACCCCC TGCACACTTCACCCCC TGCACACTTCACACCCCC TGCACACTTCACACCCCCCCCCC	21 TACCTATCAT CCAGCTATTG CAGCCACTG GTCCCCAGTG GTCCCCAGTG ATGGCAGCAC CCTCTAACT CACCTGTGT GGCTATAGA GTTTATGAAC GATGCCTGTT AACTCAGCCTGTT AACTCAGCC TGGCAGTTA ACTCAAGCC CCAGGCATTG CAAAAAGTGA ATGAACTGC CCAGGCATTA TACAGACTGC TCCTGCAGC AGGCATTA CAAAAAGTGA ATGAACAGTA TACAGACTGC CCAGGCATTA TACAGACTGC CCAGGCATTA TACAGACTGC CCAGGCATTTC CTGCAGG AAGACAATA TACAGACCAG AAGACAATA TACAGACCAG AAGACTACT CTGCTTGTGG AGATCTTCCT CTCAGCTGCT CTCAGCTGCT CTCAGCTGCT GGCACCTGCA GGGACTTTGC GGACCAGCCCC	31 TACTCGATGC GACCTTACTA TGGTCCCCAC ACGCCCCAGA GCGCCCCAGA GGACCTTCCT AGTGCTCCAA GGTGGACCAAA GCCAAGACGA ATAATTTTTA TGAACACAGA GCCAGAGCCAG GCCAGAGCCAG GCCAGAGCCG GCCAGCCCAG GCAGCTGCCC TGATGCTGCAGC GCAGTGACCC GCAGTGACCC TGCAGCC GCAGTGACCC TGCAGCC GCAGTGACCC TGCAGCC GCAGTGACCC GCACTGCCG GCACTGCCG GCACTGCCG GCACTGCCC TGATGCCC TGATGGCTC TGATGGCTC TGCCCAGCC GCACTGCCC GCACTGCCC TGCATGCCC TGCCCAGCC TGATGCCTC TGATGGCTC TGATGGCTTC	41 TGTTGATAAC TGAAAACCAT TGTTGATAAC TGAAAACCAT TGTCTACGAG GGTCCTGACG CGTGGGAGCT CTCTGGGATA CGTGTCACAC CTTCATCCTT CTGGAACGAG CTCTAGCCAA AGTGGTTTCT GATCGTGGC CAATGGTGC CAATTATCAC GACTATCAC GACTATCAC GACTATCAC GACTATCAC GACTATCAC GACTATCAC GATAGGACAT CTATGACAAC CTATGCCAG GATAGGGGT GAATGTTTT TGTCTGGCTT TGCTCTCCTT TCAATTCTGC TCAATGTCTCC TCAATGTCCAG TCAATGTCC TCAATGTCC TCAATGTCC TCAATGTCC TCAATGTCC TCAATGTCG TTCCAGGGGC AGATGAAAAA TCTGGGGCCT TTGCCTT TTCCAGGGGC TTCAAGGCCTT TTGCACTGTT TGCTTGGCCTT TCCAGGGCC TTCAAGGCCTT TTGCACTGGT TTCAAGGCCTT TTGCACTGGT TTGCACCTT TTGCACTGGT TTGCACTGGT TTGCACTGGT TTGCACTGGT TTGCACTGGT TTGCACTGGT TTGCACCTT TTGCACTGGT TTGCACCTT TTGCACCT TTGCAC	51 AGCAAGATGG GGATACCAAC GTGCATCCGG CAGGCTTCCAG CAGGCTGCCG GAGTGCCAGT TGCCCCGGCG GGATACTA ACTACGGGC GGAATACTG GTGAGATCT TTACGCTGTT GGTGAGAGCG CACATGTTCT TCAAGACCA GACTACTGT TCCTAGTTTT GCTGCAAGG CTGATCACG GTGACACG TCCTAGTCACG TCCTTGACGATTT GCTGCCAAGG CTGATCACG TCCTTGACGT ACCTTGACGATTT GCTGCCAAGG CTGATCACG TCCTTGACGT ACCCCTTGACGT ACCCCTTGACGT CACACGACGC CAATTTTGGA GGGAGACAT CTTGGTAGTG ACCCCTTGT AGGGAGACAT CTTGGTAGTG AGGAGACAT CTTGGTAGTG AGCACCCTTG AGGAGACAT CTTGGTAGTG AGCACCCTTG AGGAGACAT CTTGGTAGTG AGCACCCTTG AGGAGACAT CTTGGTAGTG AGCACCCTTG AGGACACCTTG AGGACACCTTG AGTCACCTTGT AGGTACCCTTGT AGGTACCTTGT AGGTACCTTGT AGGTACCTTGT AGGTACTTCT AGGTACCTTGT AGGTACCTTGT AGGTACCTTGT AGGTACCTTGT	60 120 180 240 360 420 480 540 660 720 1020 1020 1020 1140 1260 1320 1320 1560 1560 1680 1740 1800 1920 1980

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WO 02/30268

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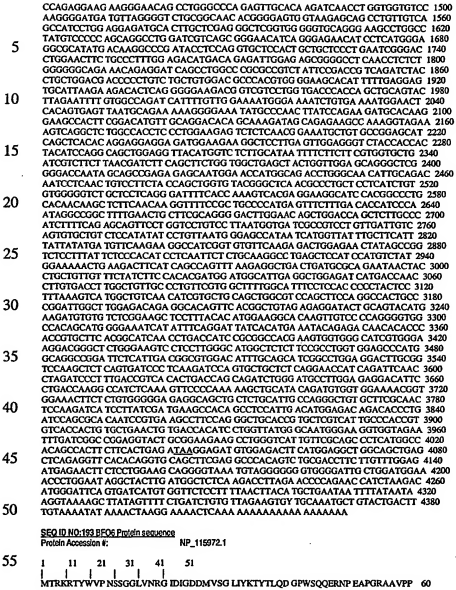
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45		3CO2 DNA sequer	10e AJ400877				oond to start and stop co	. dons)
45	SEQ ID NO:188 E	3CO2 DNA sequer	1 <u>ce</u>				pond to start and stop co	dons)
	SEQ ID NO:188 E Nucleic Acid Acce	3CO2 DNA sequer ession #: Coding	10e AJ400877				pond to start and stop co	dons)
45 50	SEQ ID NO:188 E Nucleic Acid Acce	3CO2 DNA sequer ession #: Coding	n <u>ce</u> AJ400877 sequence:				pond to start and stop co	dons)
	SEQ ID NO:188 E Nucleic Acid Acce	3CO2 DNA sequer ession #: Coding 21 31 4	nce AJ400877 sequence: 41 51	6	1-3080 (underlined			dons)
	SEQ ID NO:188 E Nucleic Acid Acce	3CO2 DNA sequer ession #: Coding 21 31 4	AJ400877 Sequence: 41 51	ECG CCGCCAC	1-3080 (underlined	d sequences corres	CTGC 60	dons)
	SEQ ID NO:188 E Nucleic Acid Acce 1 11 2	3CO2 DNA sequer ession #: Coding 21 31 4 1 1 G CACACCTCC	AJ400877 sequence: 41 51 CCCGCGCCGC	ECG CCGCCAC	1-3080 (underlined CGC CCGCACT CAA CCGTCCC	d sequences corres	CTGC 60	dons)
50	SEQ ID NO:188 E Nucleic Acid Acce 1 11 2 GGCGTCCGCC CCGCAACCGC CGGTGCTGCT	3CO2 DNA sequer ession #: Coding 21 31 4 GCACACCTCC TGAGCCATC	AJ400877 sequence: 41 51 CC CGCGCCGC	EG CCGCCAC CG CGGCCG GC CACTGCTG	1-3080 (underlined CGC CCGCACT CAA CCGTCC CT GCTGGCG	d sequences corres FCCG CCGCCTC CGGG GCGGCCC	CTGC 60 CTGGG 120 CGC 180	dons)
	SEQ ID NO:188 E Nucleic Acid Acce 1 11 2 GGCGTCCGCC CCGCAACCGC CGGTCCGGCC CGGGTCCGGGCCCGGGCCGGG	GCO2 DNA sequer ession #: Coding 21 31 4 G CACACCTCC C TGAGCCATC G CCGTGCTGCT G CCGTGCCGC	AJ400877 Sequence: 41 51 CC GCGCCGCCCCCCCGCCCCCCCCCCCCCCCCCCCCC	ECG CCGCCAC TCG CCGGCCG GC CACTGCTG AGG AGGATGT	1-3080 (underlined CGC CCGCACT CAA CCGTCCC ICT GCTGGCG TAGA TGAGTC	d sequences corres FCCG CCGCCTC CGGG GCGGCC GGG GCCGTCC	CTGC 60 CTGGG 120 CGC 180 GCTAG 240	dons)
50	SEQ ID NO:188 E Nucleic Acid Acce 1 11 2 GGCGTCCGCC CCGCAACCGC CCGGTCCGGCC CCGGTCCGGCAACCGC ATGACTGCCA	3CO2 DNA sequer ession #: Coding 21 31 4 G CACACCTCC C TGAGCCATC G CCGTGCTGCT G CCGTGCCGAC A TGCCGACG	AJ400877 sequence: 41 51	ECG CCGCCAC TCG CGGGCCG GC CACTGCTG AGG AGGATGT GA ACACACC	1-3080 (underlined CGC CCGCACT CAA CCGTCC CCT GCTGGCG CAGA TGAGTC CAC CTCCTAC	d sequences corres reced ecedeere egge decedeer regee caace race caace caac recere	CTGC 60 CTGGG 120 CGC 180 GCTAG 240 CGCA 300	dons)
50	SEQ ID NO:188 E Nucleic Acid Acce 1 11 2 GGCGTCCGCC CCGCAACCGC CGGTGCTGCT CGGGTCCGGCA ATGACTGCCA AGCCTGGCTA	3CO2 DNA sequer sssion #: Coding 21 31 4 G CACACCTCC C TGAGCCATC T GCTGCTGCT A TGCCGACGC A CCAAGGGGA	AJ400877 sequence: 41 51 CC CGCGCCGC CC ATGGGGGT GCTGCTGCCCCCCCCCCCCCCCCCCCCCCCC	CG CCGCCAC CG CGGGCCG GC CACTGCTG GA AGAACC GA ACACACC AGT GTGAGG	1-3080 (underlined CGC CCGCACT CAA CCGTCCO ICT GCTGGCG TAGA TGAGTO CAC CTCCTAC ACAT CGATGA	d sequences corres FCCG CCGCCTC CGGG GCGGCC GGG GCCGTCC FTGCC CAAGGG	CTGC 60 CTGGG 120 CGC 180 GCTAG 240 GCA 300 TGAGC 360	dons)
50	SEQ ID NO:188 E Nucleic Acid Acce 1 11 2	GCO2 DNA sequer ession #: Coding 21 31 4 G CACACCTCC C TGAGCCATC G CCGTGCCG A CCCAGGGG G CTGTGTCCA	AJ400877 sequence: 41 51 CC CGCGCCGC CC ATGGGGGT G CTGCTGCCC CG GGGCCGCC AA GGCAGGC AT GACTGTTTC	EG CCGCCAC TCG CGGGCCG GC CACTGCTG AGG AGGATGT GA ACACAC AGT GTGAGG GA ATATTCCA	1-3080 (underlined CGC CCGCACT CAA CCGTCCO CT GCTGGCGO TAGA TGAGTG CAC CTCCTAC ACAT CGATGA LGG CAATTAT	d sequences corres FCCG CCGCCTC CGGG GCGGCC GGG GCCGTCC FTGCC CAAGG AAG TGCTCCT AATGT GGAAA' CGT TGCACTT	CTGC 60 CTGGG 120 CGC 180 GCTAG 240 TGCA 300 TGAGC 360 GTT 420	dons)
50 55	SEQ ID NO:188 E Nucleic Acid Acce 1 11 2 GGCGTCCGCC CCGCAACCGC CGGTCCGGC AGCCTGGCC AGCCTGGCC AGCCTGGCC ATGACTGCCA AGCTGGCCT TCAATGGAGT TTGATGGCTT	GCO2 DNA sequer ession #: Coding 21 31 4 G CACACCTCC C TGAGCCATT G CCGTGCTGCT G CCGTGCCGC A TGCCGACGC A TGCCGACGC C CAAGGGGA C CTGTGTCCA	AJ400877 Sequence: 41 51 CC CGCGCCGC CC ATGGGGGT CC CTGTGCCC CC GGGCCGCAC CC CTGTGTCA AA GGCAGGCAT T GACTGTTTT T CATGACGGT	CG CCGCCAC CCG CCGCCCG GC CACTGCTG AGG AGGATGT GA ACACACC GA ATATTCCA TC ATAATTGT	1-3080 (underlined CGC CCGCACT CAA CCGTCCC ICT GCTGGCG TAGA TGAGTG CAC CTCCTAC ACAT CGATGA IGG CAATTAT CT TGATGTGC	d sequences corres rCCG CCGCCTC CGGG GCGGCC GGG GCCGTCC ITGCC CAAGGG AAG TGCTCCI AATGT GGAACTT GACTGCACTT GACTGCACTT	CTGC 60 CTGGG 120 CGC 180 GCTAG 240 GCA 300 TGAGC 360 GTT 420 TGG 480	dons)
50 55	SEQ ID NO:188 E Nucleic Acid Acce 1 11 2 GGCGTCCGCC CCGCAACCGC CCGCACGCC CCGGTCCGCC CCGGTCCGCC CCGCTCCGCT	3CO2 DNA sequer ession #: Coding 21 31 4 G CACACCTCC C TGAGCCATC G CCGTGCTGCT G CCGTGCCGA A TGCCGACGC A CCAAGGGGA C CATGTTGGC G CGGCTGCCG G CGGCTGCCG	AJ400877 Sequence: 41 51	CG CCGCCAC CG CGGCCG GC CACTGCTG AGG AGGATGT GA ACACAC AGT GTGAGG ATATTCG TC ATAATTGT FTG TCAACGT	1-3080 (underlined CAA CCGTCC CAA CCGTCC CT GCTGGCG CAGA TGAGTC CAC CTCCTAC ACAT CGATGA GG CAATTAG CAT GGGGAG	d sequences corres CCCG CCGCCTC CGGG GCCGCCC FTGCC CAAGGG AAG TGCTCCT AATGT GGAAATT GAC GAGTGCC CTAT GAGTGCC	CTGC 60 TTGGG 120 CCGC 180 GCTAG 240 TGCA 300 TGAGC 360 GTT 420 TGG 480 TGCT 540	dons)
50	SEQ ID NO:188 E Nucleic Acid Acce 1 11 2 GGCGTCCGCC CCGCAACCGC CGGTGCTGCT CGGGTCCGCC AGCCTGGCT TCAATGGAG TTGATGGCT AGAACAATG GCAAGGAGG	3CO2 DNA sequer sission #: Coding 21 31 4 G CACACCTCC C TGAGCCATC G CGTGCCGC A TGCCGACGC A CCAAGGGG C CTGTGTCCA C CATGTTCGC G CGGCTGCCC G CGGCTGCCC G GTTTTTCCT	AJ400877 Sequence: 41 51	EG CCGCCAC CG CGGCCG GC CACTGCTG GA ACACACC AGT GTGAGG, GA ATATTCCA IC ATAATTG ITG TCAACGT ITC AGCACAC	1-3080 (underlined CGC CCGCACT CAA CCGTCCO CT GCTGGCG CAC CTCCTAC ACAT CGATGA GG CAATTAT CT TGATGTG CAT GGGGAG CTG CATTCAC	d sequences corres CCCG CCGCCTC CGGG GCCGCC CGG GCCGCCC AGGG AAG TGCTCCT AATGT GGAAA CGT TGCACTT GACTGC CTAT GAGTGC CCCTAT GAGTGC	CTGC 60 CTGGG 120 CGC 180 GCTAG 240 GCA 300 TGAGC 360 GTT 420 TGG 480 TGCT 540 GAGG 600	dons)
50 55	SEQ ID NO:188 E Nucleic Acid Acce 1 11 2 GGCGTCCGCC CCGCAACCGC CGGTGCTGCC AGCCTGGCTA TCAATGGAG TTGATGGCTA AGACAATG GCAAGGAGG GCAAGGAGG GCCTGAGCTC	GCO2 DNA sequer session #: Coding 21 31 4 G CACACCTCC C TGAGCCATCC C TGCTGCTGCT A TGCCGACGC A CCAAGGGGA C CATGTTGCC G CGGCTGCCA G CGTTTTCCT G CATGAATAA	AJ400877 Sequence: 41 51 CC CGCGCCGC CC ATGGGGGT G CTGCTGCC CC CTGTGTCA AA GGCAGGC AT GACTGTTT T CATGACGT G AGTGACAA AG GATACCGT	ECG CCGCCAC CCG CCGGCCG GC CACTGCTG GA ACACACC AGT GTGAGG, GA ATATTCCA TC ATAATTGT STG TCAACGT TC AGCACAC	1-3080 (underlined CGC CCGCACT CAA CCGTCCG CT GCTGGCGG TAGA TGAGT CAC CTCCTAC ACAT CGATGA GG CAATTAT CT TGATGTGG CAT GGGGAG CTG CATTCAC	d sequences corres CCCG CCGCCTC CGGG GCCGCCC GGG GCCGTCCC AAG TGCTCCC AATGT GGAAA' CGT TGCACTT GAC GAGTGCC CTAT GAGTGCC CCTAT GAGTGCC CCGC TCGGAAA'	CTGC 60 CTGGG 120 CGC 180 GCTAG 240 GCA 300 TGAGC 360 GTT 420 TGG 480 TGGT 540 GAGG 660	dons)
50 55	SEQ ID NO:188 E Nucleic Acid Acce 1 11 2 GGCGTCCGCC CCGCAACCGG CGGTGCTGCC CGGTCCGGG ATGACTGCCA AGCCTGGCT AGACCATGGAG TTGATGGCT AGACAATGG GCAAGGAGGG GCCTGAGCT GCAAGCAGG	GCO2 DNA sequer ession #: Coding 1 31 4 G CACACCTCC C TGAGCCATC G CCGTGCCGC A TGCCGACGC A TGCCGACGC C CAAGGGGC C CATGTTGCC G CGTTTTTCC G CGTTTTTTCC G CGTTGAGTCC C TGAGAATAA C CTGTGAGTC	AJ400877 Sequence: 41 51 CC CGCGCCGC CC ATGGGGGC CC GGGCCGCC CC GGGCCGCC CC CTGTGTCA AA GGCAGGC TT CACTGTTTT T CATGACGGT AG CATACCTC TG AGTGACAA AG GATCACGC CC AGGCCTGG	CG CCGCCAC TCG CGGCCG GC CACTGCTG AGG AGGATGT GA ACACAC GC ATAATTGT GT GTCAACGT TC AGCACAC GCT GTAGTCA	1-3080 (underlined CAA CCGTCCC CAA CCGTCCC CAG TGAGGG CAC CTCCTAC ACAT CGATGA GG CAATTAT CT TGATGTGC CAT GGGGAG CTG CATTCAAC GGC CAAGAAC	d sequences corres FCCG CCGCCTC GGG GCGGCC FTGCC CAAGGC AAG TGCTCCI AATG TGCACTT GAC GAGTGCC CTAT GAGTGCC CGC TCCGCAA CCAG AGAGAC	CTGC 60 CTGGG 120 CGC 180 GCTAG 240 TGCA 300 TGAGC 360 GTT 420 TGG 480 TGCT 540 GAGG 600 AGGG 660	dons)
50 55 60	SEQ ID NO:188 E Nucleic Acid Acce 1 11 2 GGCGTCCGCC CCGCAACCGC CCGGTCCGCT CCGGTCGCT TCAATGGACT AGACAATGG GCAAGGAGG GCCTGAGCT GCAGGCTCGCT TCTGACCTGCT TCTTGACCTG	CO2 DNA sequer ession #: Coding 21 31 4 G CACACCTCC C TGAGCCATC G CCGTGCTGCT G CCGTGCTGCT G CCGTGCTGCT G CTGTTTCCT G CGGCTGCCG G GTTTTTCCT G CGGCTGCAC C CATGAGAC C TATGAGAC C TATGAGAC C TTGTGAGT G CTGTGAGT G CTGTGAGT G CTGTGAGT G CTGTGAGT G TAACCATGG	AJ400877 Sequence: 41 51 CC CGCGCCGC CC GGGGCCGC CC GGGCCCGC CC CTGTGCA AA GGCAGGC AT GACTGTTA T CATGACGG AG CATACCTC G AGTGACAA GCATGCACA AG CATACCTC G AGTGACAA AG CATACCTC G AGTGACAA AG CATACCTC AG AGTGACAA AG CATACCTC	CG CCGCCAC CG CGGCCG GC CACTGCTG AGG AGGATGT GA ACACACC AGT GTGAGG GC ATAATTGT TG TCAACGT TC AGCACAC TC TCAGCACAC GT GTAGTCA GT GTAGTCA	1-3080 (underlined CAA CCGTCCC CAA CCGTCCC CAGA TGAGTC CAC CTCCTAC ACAT CGATGA CGG CAATTAA CGT GAGGAG CTG CATTCAC CAT CTGCAA CAT CTGCAA CAT CTGCAA	d sequences corres CCCG CCGCCTC CCGG GCCGCCC FTGCC CAAGGG CAAG TGCTCCT AATGT GGAAAT GAC GAGTGCC CTAT GAGTGCC CTAT GAGTGCC CTAT GAGTGCC CTAT GAGTGCC CCAGGAAAGGCCC CCAGGAAAGGCCC CCAGGAAAGGCCCACACAGACACACAC	CTGC 60 TTGGG 120 CCGC 180 GCTAG 240 TGCA 300 TGAGC 360 GTT 420 TGG 480 TGCT 540 GAGG 600 AGGG 660 TGCA 720 GATG 780	dons)
50 55 60	SEQ ID NO:188 E Nucleic Acid Acce 1 11 2 GCGTCCGCC CCGCAACCGC CCGCACCGC CGGTGCTGCT CGGGTCGCC AGCCTGGCT TCAATGGAGC TTGATGGCT AGAACAATG GCAAGGAGG GCCTGAGCTC GCAACCGCC GCCCAGAGTC GCCCAGAGTC GCCCAGAGTC	3CO2 DNA sequer sission #: Coding 21 31 4 G CACACCTCC C TGAGCCATC G CCTGCTGCCG A TGCCGACGC A CCAAGGGG C CTGTGTCCA G CGGCTGCCC G GGTTTTCCT G CATGAATAA C CTGTGGACG C TTGTGGCG G TAACCATGG G TAACCATGG G CAGCTGCCA	AJ400877 Sequence: 41 51	CG CCGCCAC CG CGGCCG GC CACTGCTG GA ACACACC AGT GTGAGG, GA ATATTCCA GC ATAATTGT TC ATAATTGT TC AGCACAC GCT GTAGTCA ATT TTGAGCT GT GCCAGCA CA AGATGCA	1-3080 (underlined CGC CCGCACT CAA CCGTCCC CAG TGCTGCCG CAC CTCCTAC ACAT CGATGA GG CAATTAT CAT GGGGAG CTG CATTCAC CAT CTGCAAC GGC CAAGAAC CTC CTGTGAAC	d sequences corres CCCG CCGCCTC CGGG GCCGCC CGGG GCCGCC CAAGG CAAG TGCTCCT ATGT GGAAA CGT TGCACTT CATGT GAGTGCC CTAT GAGTGCC CTAT GAGTGCC CCGC TCGGAAG CGAG GCCCCA CGAG AGAGAG CGAT ACAGCCC GAGAG AGCACC GAGAG AGCACC CGAT ACAGCCC CGAG AGCACC CGAT ACAGCCC CGAG AGCACC CGAGAG AGCACC CGAGAG AGCCCC	CTGC 60 CTGGG 120 CGC 180 GCTAG 240 GCA 300 TGAGC 360 GTT 420 TGG 480 TGGT 540 GAGG 660 TTGCA 720 GATG 780 CCTTG 840	dons)
50 55	SEQ ID NO:188 E Nucleic Acid Acce 1 11 2	GCO2 DNA sequer ession #: Coding 21 31 4 G CACACCTCC G TGAGCCATC G CCGTGCCGC A TGCCGACGC G CGTGTGCCA G CGAGGGGA G CTGTTGTCCA G CATGTTGCC G CAGCTGCCA G CATGTTGCC G CAGCTGCCA G CATGAATAA C CTGTGAGTG G TAACCATGC G CAGCTGCCA A CACTGTCCCA	AJ400877 Sequence: 41 51	CG CCGCCAC CG CGGCCAC GC CACTGCTG AGG AGGATGT AGG ACACCC GC ATAATTGT TTC ATAACTT TTC AGCACAC GCT GTAGTCA TTTTTGAGCTT GT GCCAGCAC CA AGATGCA CAG AGAGCAC	1-3080 (underlined CGC CCGCACTI CAA CCGTCCC CAC CTCCTAC CAC TCCATAC ACAT CGATGA GG CAATTAT CT TGATGTGC CAT GGGAGG CTG CATTCAC CAT CTGCAAC GGC CAAGAAC CTC CTGTGAAC	CCG CCGCCTC CGGG GCGCCC GGG GCGCCC GGG GCGCCC AGG GCCGCC AGG GCGCC CAAGG	CTGC 60 CTGGG 120 CGC 180 GCA 300 TGAGC 360 GTT 420 TGGT 540 GAGG 660 AGGG 660 TGCA 720 GATG 780 CCTTG 840 TGGGG 900	dons)
50 55 60	SEQ ID NO:188 E Nucleic Acid Acce 1 11 2 GCGGTCCGCC CCGCAACCGC CCGGTCCGCCA AGCCTGCCCA AGCCTGCCT TCAATGAGG TTGATGGCTT AGAACAATGG GCAAGGAGGG GCTGAGCTCGC TCTTGACCTG GCCCAGAGTCG AGCAGAGGG ATAAACCGGG ATAAACCGGG	GCO2 DNA sequer ession #: Coding 21 31 4 G CACACCTCC C TGAGCCATC G CTGTGCTGCT G CCGTGCCG A TGCCGACGC A TGCCGACGC C CAAGGGGA C CAAGGGGA C CATGTTGCC G CTTTTTCCT G CATGAATAA C CTGTGAGTC G TAACCATGG G TAACCATGG G A CACTGCC	AJ400877 Sequence: 41 51	CG CCGCCAC CG CGGCCG CG CGGCCG CG AGGATGT GA ACACAC CG ATATTCCA TC ATAATTGT GT GTCAACGT TTC AGCACAC TTTTTGAGCA TTTTTGAGCT TTTTGAGCT TTTTTGAGCT TTTTTGAGCT TTTTTGAGCT TTTTTGAGCT TTTTTGAGCT TTTTTTGAGCT TTTTTTGAGCT TTTTTTGAGCT TTTTTTTTTT	1-3080 (underlined CAA CCGTCCC CAA CCGTCCC CACAT GATGAC CACAT CGATGA CAT GGGGAG CTG CATTCAC CAT GGGGAG CTG CATTCAC CAT CTGCAAC CAC CTGTGAC CAC CTGTGAC CAC AGATGA CTC CTGTGAC	d sequences corres FCCG CCGCCTC CGGG GCGGCC FTGCC CAAGGG AAG TGCTCCT AATGT GGAAAT GGACTT GAC GAGTGCC CTAT GAGTGCC CTAT GAGTGC CTAT GAGTGC CGC TCGGAG CGAG GCCCAA CCAG AGAGAC CGAG AGCTG CAGT GGAGAC CAGT GGGGAC CAGT GGAGGC	CTGC 60 CTGGG 120 CCGC 180 GCTAG 240 GCA 300 TGAGC 360 GTT 420 TGG 480 TGCT 540 GAGG 660 TGCA 720 GATG 780 CCTTG 840 TGGGG 900 CTGTG 960	dons)
50 55 60	SEQ ID NO:188 E Nucleic Acid Acce 1 11 2 GCGGTCCGCC CCGCAACCGC CCGGTCCGCT CCGGTCGGCT TCAATGGACT AGACAATGG GCAAGGAGG GCTTGACTG GCAGGTCGC TCTTGACCTG GCACGACGT AGACAATGG AGCGTCGG AGCACGTCG AGCACGTCG AGCACGTCG AGCACGTCG ACCACACTC	BCO2 DNA sequer Ession #: Coding 21 31 4 G CACACCTCC C TGAGCCATC G CTGCTGCTGCT G CCGTGCTGCT G CCGTGCTGCT G CCGTGCTGCT G CTGTTTCCT G CGGCTGCCC G GTTTTTCCT G CTGTGAGC G CATGATACA C CTGTGAGT G CATGATACA C CTGTGAGT G CACATGCC A CACTGCCC A CACTGCCC A TGAAACGGC G TAAGGATAC	AJ400877 Sequence: 41 51 CC CGCGCCGC CC ATGGGGGT CC CTGCTGCCC CC GGGCCCGC CC CTGTGCA AA GGCAGGC AC CTGTGCA AA GGCAGGC CC CTGTGCA AA GGCAGGC AC CATACCTC CG AGTGACAA AG CATACCTC CG AGGCTGG CG AACGGTGG AT CCACAGTAA CT CCACAGTAA CT TCGACAGG	CG CCGCCAC CG CGGCCGC GG CGGGCCG GG AGGATGT GA ACACACC AGT GTGAGGA TC ATAATTGT TG TCAACGT TC TCAGCACAC GT GTAGTCA GT GCCAGCA CA AGATGCA CAG AGAGCA TG TCCACTGC	1-3080 (underlined CAA CCGCACT CAA CCGTCCC CAG TGAGGG CAG CTCCTAC ACAT CGATGA CGT CAATTAA CGT CAATTAA CGT CATTCAC CAT CGCAAG CGC CAAGAA CTC CTGCAA CAC ACAGT CTC CTGTGAC CAC CACAT CTGC CACTAG CAC CACAT CTGC CACTAG CTGC CACTAG CTGC CACTAG CTGC CACTAG CAG TTGTCCT	CCG CCGCCTC CGGG GCGCCC TGCC CAAGGG CAAG TGCTCCT ATGT GGAAAT GAC GAGTGCC CTAT GAGTGCC CAG AGAGAC CAGA GCCCCA CCAG AGAGAC CAAT GGAGGC CAAT GGAGGG GTT GGATTCA	CTGC 60 CTGGG 120 CGC 180 CGC 180 CGCA 300 TGAGC 360 CGTT 420 TGG 480 TGGT 540 GAGG 600 AAGGG 660 TGGA 720 GATG 780 CCTTG 840 TGGGG 900 CTGTG 960 CTC 1020	dons)
50 55 60	SEQ ID NO:188 E Nucleic Acid Acce 1 11 2 GGCGTCCGCC CCGCAACCGC CGGAACCGC CGGTGCTGCT TCAATGGAGG TTGATGGCT AGAACAATG GCAAGGAGG GCCTGAGCTC GCAGCTCG TCTTACCTG GCAGCTCG AGCAGCTCG TCTTACCTG TCTTACCTG TCTTACCTG TCTTACCTG TCTTACCTG TCTTACCTG TCTCAGTTGACTT TCAGTTGACTT	3CO2 DNA sequer session #: Coding 21 31 4 G CACACCTCC C TGAGCCATC G CGTGCTGCT G CCGTGCTGCT G CCGTGCTGCG A TGCCGACGG A TCCAGGTGCCA G CTGTTTCCT G CATGTTCCA G CAGCTGCCA G CTGTTTCCT G CATGAATAA C CTGTGAGC G TAACCATGG G CAGCTGCCA A CACTGTCCA A TGAAACGGC G TAAACGGC G TAAAGGAAAACGG T GAAACGGC G TAAGGAAAAA	AJ400877 Sequence: 41 51 CC CGCGCCGC CC ATGGGGGT GC CTGCTGCCC CC GGGGCCGCA CC CTGTGTCA AA GGCAGGCA AA GGCAGGCA AG CATACCTC AG AGTGACAA AG GATACCTC AG AGTGACAA AG GATCACGC AG CATACCTC AG CATACCTC AG AGTGACAA AG GATCACGC AG CATACCTC AG AGTGACAA AG GATCACGC AG CATACCTC AG AGTGACAA AG CATACCTC AG AGTGACAA AG CATACCTC AG AGTGACAA AG CATACAGGC AT CCACAGTA AG CATGACAA AG CATACCTC AG AG CATACCT	CG CCGCCAC CG CGGCCG CG CGGCCG AGG AGGATG GA ACACAC AGT GTGAGG, GA ATATTCCA TC ATAATTGT TTC AGCACAC GT GTAGTCA AGT GCAGCA CA AGATGCA CA AGATGCA CGG AAACGTA TGG TCCACTGG TTG TCCACTGG	1-3080 (underlined CAA CCGCACT CAA CCGTCCC CT GCTGGCG CAC CTCCTAC ACAT CGATGA GG CAATTAT CT TGATGTGC CAT CGGGAG CTG CATTCAC CAT CTGCAAC CTC CTGTGAC CAC AGATGG ACAC CACATC CTGCTGAC CAC TGTGAC CAC TGTGAC CAC TGTGAC CAC TGTGAC	d sequences corres CCCG CCGCCTC CGGG GCCGCCC EGGG GCCGCCC EGCC CAAGGC CAAG TGCTCCT ATGT GGAAA CGT TGCACTT GACTGC CTAT GAGTGCC CTAT GAGTGCC CTAT GAGTGCC CGC TCGGAAC CGAT ACAGCCC GAGG AGCCCCA CGAT GCAGGC CCAT GGATCA CCAT GGATCA CCAT GGATCA CCGC AATGGA CCGC AATGGA	CTGC 60 CTGGG 120 CGC 180 GCTAG 240 GCA 300 TGAGC 360 GTT 420 TGG 480 TGGC 540 GAGG 660 TGGA 720 GATG 780 CCTTG 840 CCTTG 840 CTGGG 900 CTTGTG 960 CTC 1020 GGTT 1080	dons)
50556065	SEQ ID NO:188 E Nucleic Acid Acce 1 11 2	BCO2 DNA sequer ssion #: Coding 21 31 4 G CACACCTCC G TGAGCCATC G CCGTGCCGC A TGCCGACGC G CGTGCCCG G CAGGGGACGC G CATGTTGCCA G CATGTTGCC G CATGTTGCC G CATGTTGCC G CATGTTCC G CATGTTCC G CATGAATAA C CTGTGAGT G CAGCTGCC A TGCGAAGAC C TGAAACGCC A TGGGAAGAC C TGGAAAAA	AJ400877 Sequence: 41 51	CG CCGCCAC CG CGGCCAC GC CACTGCTG AGG AGGATGT AGG ACACCC GC ATAATTGT TC ATAACTT TC AGCACAC GCT GTAGTCA GT TTGAGCTC AGCAGCA CA GATGCA CA GAGAGCA CA GAGAGCA CA GATGCA CA GATTTGATGA CA GTTTTGATGA	1-3080 (underlined CGC CCGCACTI CAA CCGTCCT CCT GCTGGCGG TAGA TGAGTG CAC TCCTAC GG CAATTATT CT TGATGTGG CAT GGGGAGG CTG CATTCAC CAT CTGCAAC GGC CAAGAAC CTC CTGTGAAC GGC CAGATGG CAC AGATGG CAC AGATGG CAC TGTCAAC CAG TGTCCAC TGC TGTCAAC CAG TTGTCCAC TGG CCAGACC	CCG CCGCCTC CGGG CCGCCCC GGG GCGCCC GGG GCGCCC CAAGGC CAAGGC CAGG TGCACTT GAAAA CGT TGCACTT GACT CGC CTGGAA CGG GCCCA CGAG AGAGAC CGAG AGAGAC CAGT GCACT CAGT GATGAC CAGT GATGAC CAGT GATGAC CAGT GATGAC CAGT GATGAC CACT	CTGC 60 CTGGG 120 CGC 180 GCTAG 240 GCA 300 TGAGC 360 GTT 420 TGG 480 TGGT 540 GAGG 660 AGGG 660 TGCA 720 GATG 780 CCTIG 840 TGGGG 900 TTGTG 960 CTC 1020 GGTT 1080 TTTA 1140	dons)
50 55 60	SEQ ID NO:188 E Nucleic Acid Acce 1 11 2 GCGCTCCGCC CCGCAACCGC CCGCAACCGC ATGACTGCCA ATGACTGCCA ATGACTGCCT AGAACAATGG GCAAGGAGGG GCCTGAGCTC GCACCAGAGTC GCACCAGAGTC AGCAGAGTC AGCAGAGTC ACCAGAGTC ACCAGAGTC ACCAGAGTC ACCACTTTGACCTT ACCACTTTGACCTT ACCACTTTGACCTT ACCACTTTTTACCTT ACTACTTTTACCTT ACTACTTTTACCTT ACTACTTTTACCTT ACTACTTTTACCTT ACTACTTTTACCTT	GCO2 DNA sequer ression #: Coding 21 31 4 G CACACCTCC C TGAGCCATC G CTGTGCTGCT G CCGTGCCG A TGCCGACGC A TGCCGACGC C CAAGGGGA C CAAGGGGA C CATGTTGCC G CTTTTTCCT G CATGAATAA C CTGTGAGTC G TAACCATGG G TAACCATGG G TAACCATGG G TAACCATGC T GAAACGCC T GAAACGCC T GAAACGCC T GAAACACC C TGCAAAAA C CTGCAAAAA C CTGCAAAAA C AGATGAGAA	AJ400877 Sequence: 41 51 CC CGCGCCGC CC ATGGGGGT CC CTGCTGCCC CC GGGCCGCC CC CTGTGTCA AA GGCAGGGT T CATGACGGT AG CATACCT CG AGGCTGG GACCAGG GACCAGG CT CACAGGT AT CACAGGT CT CACAGGT CT CACAGGT CT CACAGGT CT CACAGGT CT CACAGGT CT CCACAGGT CT TCGACAGG CA TGTAAAGGA CA TTGTGGGG AG TCTTGCCA	CG CCGCCAC CG CGGCCG CG CGGCCG CG AGGATGT GA ACACAC CG ATATTCCA TC ATAATTGT GT TCAACGT TTG TCAACGT TTG TCAACGT TT TTGAGCA TT TTGAGCA CA AGATGCA AGATGCA TG TCACTG TG ATTTGAG AG	1-3080 (underlined CAA CCGTCCC CAA CCGTCCC CACAT GATGAC CACAT CGATGA CAT GATGAC CAT GGGGAG CTG CATTCAC CAT CTGCAAC CAC CTGTGAC CAC CAGATGAC CAC CACATGAC CAC CACATGAC CAC CACATGAC CAC CACATGAC CAC CACATGAC CAC CACATGAC CAC TGCCAC CTGC TGTCAAC CTGC TGTCAAC CTGC TGTCAAC CTGC TGTCAAC CTGC TGTCAAC	CCG CCGCCTC CGGG GCGCCC CGGG GCGCCC CGGG GCGCCC CGTGC CAAGGC CAAG TGCTCCT CGT TGAAAT GAC GAGTGCC CTAT GAGTGCC CTAT GAGTGCC CTAT GAGTGCC CGAG GCCCCA CCAG AGGAGC CAAT GGAGCC CAAT GGAGCC CAAT GGAGCC CAT GGATCA CCAAT GGAGCC CAT GGATCA CCAAT GGAGCC CAT GGATCA CCG AATGGA CCG AATGGA CTG GATAGGA CTG GATAGGA CTG GATAGGA CTTG GATAGGA CTTG GATAGGA CTTG GATAGGA	CTGC 60 CTGGG 120 CCGC 180 GCTAG 240 GCA 300 TGAGC 360 GTT 420 TGG 480 TGCT 540 GAGG 660 AGGG 660 TGCA 720 GATG 780 CCTTG 840 CCTG 840 CTGTG 960 CTC 1020 GGTT 1080 CTTA 1140 ACCT 1200	dons)
50556065	SEQ ID NO:188 E Nucleic Acid Acce 1 11 2 GCGGTCCGCC CCGCAACCGC CCGGAACCGC CCGGTCCGGCT CCGGTCCGGCT CCGGTCGGGG ATGACTGCC AGCCTGGCT AGAACAATG GCAAGGAGG CCTGAGCT GCAGCGTCGG TCTTGACCTG GCCCAGAGTC ACCGCACCTC TCCAGTTGGA GTGATCACT TCCAGTTGGA GTGACCACAC	CO2 DNA sequer ession #: Coding 21 31 4 G CACACCTCC C TGAGCCATC G CTGCTGCTGCT G CCGTGCTGCT G CCGTGCTGCT G CCGTGCTGCT G CTGTGTCCA A TGCCGACGC A TGCCGACGC G CTTTTTCCT G CATGATCA C CTGTGAGTG G CAGCTGCCA T CAACATGCC T GAAACGGC T TGAACATGC T GAAACGGC T TGAAGAAC A TGGGAAGAC C TGCAACAA C AGATGAGAA	AJ400877 Sequence: 41 51 CC CGCGCCGC CC ATGGGGGT CC CTGCTGCCC CC GGGCCCGC CC CTGTTCA AA GGCAGGC AA GGCATGTTA T CATGACGG AG CATACCTC G AGGCCTGG G AACGGTGG AT CCACAGTA CT TCGACAGG AT CCACAGTA CT TCGACAGG CA TGCTCAT CT TCGACAGG CA TGTAAAGA CC ATCGTGGC AG CATCCTGG CA CGCCTGG CA CGCCTGG CA CGCCCTGG CA CGCCCTGG CA CGCCCTGG CA CACCCTGG CA CACCCTGC CA CACCCTGG CA CACCCTGG CA CACCCTGG CA CACCCTGC CA CACCTTGC CACCTGC CA CACCCTGC CACCCTGC CA CACCCTGC CA CACCCTCC CA	CG CCGCCAC CG CGGCCAC CG CGGGCCG GG AGGATGT GA ACACACC AGT GTGAGGA GA ATATTCCA TG TCAACGT TG TCAACGT TT TTGAGCAC GT GCCAGCA CA AGATGCA AGA TGCACAC TG TCCACTGC TG	1-3080 (underlined CAA CCGCACT CAA CCGTCCC CAG TGAGGG CAG CTCCTAC ACAT CGATGA CGG CAATTAAC CAT GGGGAG CTG CATTCAC CAT CTGCAA CAC CACAT CTGC CAGAGG CTG CTGTGAC CAC CACAT CTGC CAGAGG CTG CTGCAA CAG TTGTCCT CTG CGGCTGC TGG CGGCTGC TGG GGGCTGC TGG GGCTGCT	CCG CCGCCTC CGGG GCGCCC GGG GCCGTCC GGG GCCGTCC GTGCC CAAGG CAAG TGCTCCT ATGT GGAAAT GGT TGCACTT GAC GAGTGCC CTAT GAGTGC CCGT TGCACAC CCAG AGGAC CCAG AGGAC CAGT GCCCAC CCAG GGCCCAC CCAGT GCCCAC CCAGT GCGCC CAGT GCGCC CAGT GCGCC CAGT GCGCC CAGT GCGCC CTT GGATTCAC CCGC AATGGA AAG AAAGGA TTG GATAGGG AAC CGAGGGT	CTGC 60 CTGGG 120 CGC 180 CGC 180 CGCA 300 TGAGC 360 GTT 420 TGG 480 TGGT 540 GAGG 600 AGGG 660 TGGA 720 GATG 780 CCTTG 840 TGGA 780 CCTG 1020 GGTT 1080 TTGTA 1140 ACCT 1200 CACA 1260	dons)
50556065	SEQ ID NO:188 E Nucleic Acid Acce 1 11 2 GCGGTCCGCC CCGCAACCGC CGGAACCGC CGGTGCTGCT TCAATGGACT AGACAATGG GCAAGGAGG GCCTGAGCTC GCAGCTCG GCAGCTCG TCTTGACCTG AGCCAGAGT AGCCAGAGT AGCAGAGG ATAAACGG ATAAACGG ATAAACGG ATAAACGGC ACCGCACCTC TCCAGTTGGA GTGACCACCAC CCCTGTATGGC CCCTGTATGGC CCCCTGTATGGC	BCO2 DNA sequer Ession #: Coding 21 31 4 G CACACCTCC C TGAGCCATC G CTGCTGCTG A TGCCGACGC A TGCCGACGC A TCATGTTGGC G CATGTTCCT G CATGTTCCT G CATGTTGCC G CATGTTCCT G CATGACATAA C CTGTGCCA A TAACCATGC G TAAGCATGC G TAAGGATAC A TGGGAAGAC A TGGGAAGAC A TGGGAAGAC A TGGGAAGAC C TGCAAAAA G CTGCATCAC	AJ400877 Sequence: 41 51 CC CGCGCCGC CC AIGGGGGT GC CTGCTGCCC CC GGGGCCGCC CC CTGTGTCA AA GGCAGGC AT GACTGTTTC GG AGTGACAG AG CATACCTC GG AGTGACAA AG GATCACGC AG CATACCTC AG AGTGACAA AG GATCACGC AT CCACAGTA AT CCACAGTA AT CCACAGTA AT TCGACAGG ATCGTGGC ATCGTGGG ATCGTGGG ATCGTGGG ATCGTGGG ATCGTGGG ATCGTGGG ATCGTGGG ATCGTGGG ATCGTGGG AC CACCCTGG CTGTGGAGAG AC CACCCTGG	CG CCGCCAC CG CGGCCAC CG CGGGCCG AGG AGGATG AGG AGGATG AGG AGATATTCA TC ATAATTG TT AGCACAC AGT GTAGGCA AGT GTAGGCA AGT GTAGGCA AGATGCA AGATGTGC AGATTTGAC AGATGTGC CA CCAATGAC CA CCAATGAC	1-3080 (underlined CGC CCGCACT CAA CCGTCCC CT GCTGGCG CAG TGATGA GG CAATTAC CAT GGGGAG CTG CATTCAC CAT CTGCAAC CAT CTGCAAC CAC CACATC CAC AGATGG ACAC CACATC CTG CGCTGC CTG CGGCTGC TGG CGGCTGC TGG TGCTCT TTG TGCTTGCT TTG TGCTTGCTTG	CCCG CCGCCTC CCGG GCGCCCC CGGG GCCGCCC CAGG GCCGCCC CAGG TGCCCC CATG GGAAC CCGC TCGGAAC CGAT GCGCC CAAT GAGGCC CAAT GAGGC CTT GGATTCA CCGC AATGGA AAG AAAGGAC CACGAGAC CACGAGAC CCGC CAATGGA AAC CACGAGGC CAACGAC CACCGAGGGC CACCGAGGGC CAACGAC CCGCAACGAC CCGCAACGAC CCGCAACGAC CCGCAACGAC CCGCAACGAC CCACCGAGGGT CAACCGACGAC CCCCCCCCCC	CTGC 60 CTGGG 120 CGC 180 GCTAG 240 GCA 300 TGAGC 360 GTT 420 TGG 480 TGGC 540 GAGG 660 TGCA 720 GATG 780 CCTTG 840 CCTTG 840 CTGG 960 CTC 1020 GGTT 1080 FTTA 1140 ACCT 1200 ACCT 1200 ACCA 1260 GGCT 1320	dons)
50556065	SEQ ID NO:188 E Nucleic Acid Acce 1 11 2 GGCGTCCGCC CCGCAACCGC CCGCAACCGC CGGTCGGGCT AGACCAGCT TCAATGGAGT TGATGGCT AGACAATGG GCAAGCAGG GCCTGAGCT GCAGCTGAGCT GCAGCTGAGCT TCTTGACCTG GCAGCTGAGCT TCTGACTGAC AGCGAGAGG ATAAACGGG ATAAACGGG ATAAACGGG ATAAACGGG ATAAACGGG ATAAACGGG ATCACTTT AATTATTAAC GTGACCACACG CCCTGTATGG	GCO2 DNA sequer ression #: Coding 21 31 4 G CACACCTCC G GCGCTGCCG G CCGTGCCGC G CCGTGCCGC G CAGGGGA G CTGTGTCCA G CATGTTGCC G CAGCTGCCC G CAGGGGA G CTGTGCCC G CAGGGGA G CTGTGCCC G CAGGGGA C CATGTTGCC G CAGGATAA C CTGTGAGTC G CAGGATAA C CTGTGAGTC G CAGGAAAA C CTGCAAAAA C AGATGAGAA G CTTCACCA G CTTCACCA G CTTGCACAA C CTGTGTGAA C CTGTGTGAA C CTGCATCAA C CTGCTGTGAA C C CTGCTGTGAA C C C C C C C C C C C C C C C C C C C	AJ400877 Sequence: 41 51 CC GCGCCGC CC ATGGGGGT G CTGCTGCC CC GGGCCGC CC TGTGTCA AA GGCAGGC AT GACTGTTT T CATGACGGT AG CATACCT G GAGTGACA CG AGTCACAG CG AGCCTGG GACGCTGA AT CACAGTA CG CATCACT CT TCGACAGG AT CCACAGTA CG CATCACAG CC ATCGTGGG AG CTTTCCA ACC CTGGGG ACC CTGTGGAGA CC ATCGTGGGG ACC CTGTGGAGAC ACC CACCTGG ACTGTGGAGAC ACC CACCTGGG ACTGTGGAGAC ACC ACCTGGG ACTGTGGAGAC ACC ACCTGGG ACTGTGGAGAC ACCAGTGGGG ACTGTGGAGAC ACCAGTGGGG ACTGTGGAGAC ACCACCTGGG ACTGTGGAGAC ACCACCTGGG ACTGTGGAGAC ACCACCTGGG ACCACCTGGG ACTGTGGAGAC ACCACCTGGG ACCACCTGGG ACCACCTGGG ACCACCTGG ACCACCTG ACCACCTGG ACCACCTG ACCACCTGG ACCACCTG ACCACCTGG ACCACCTGG ACCACCTG ACCACCTG ACCACCTG ACCACCTG ACCACCTG ACCACCT ACCAC	CG CCGCCAC CCG CCGCCAC CCG CCGCCCC GC CACTGCTG AGG AGGATGT GG ACACACC GCT GTGAGGA GCA ATATTCCA TC ATAATTGT TC AGCACAC GCT GTAGTCA AGT TTGAGCT GCAGCAC AGA GCACAC AGA GCACTTTGAC AGA GCACTTTGAC AGA CATTTGAC AGA CATTTGAC AGA CATTTGAC ACA CATTTGAC CA CCAATGAC ACA CCAATGAC ACA CCAATGAC ACA CCAATGAC ACCAATGAC ACCACTGAC ACCACTCAC ACCACTGAC ACCACTCAC ACCA	1-3080 (underlined CGC CCGCACT CAA CCGTCCC CCT GCTGGCG TAGA TGAGTG CAC TCCTAC ACAT CGATGAC CAT GGGGAG CTG CATTCAC CAT CTGCAAC CAC CAGTGG CAC AGATGG ACAC CACATG TGC CAGAC TGC CAGAC TGC CAGAC TGC CAGAC TGC CAGAC TGC CGGCTGC TGC TGCTTGC TGC TGCTTGC TGC TGCTGC TGC TGCTTGC TGC TGCTTGC TGC TGCTTGC TGC TGCTTGC TGC TGCTTGC TGC TGCTTGC TGC TGCTTGC TGC TGCTTGC	CCG CCGCCTC CGG CCGCCTC CGGG GCGGCC CGGG GCGCCC CGGG GCGCCC CGG GCGCCC CAAGGC CAAG TGCACTT GACTGCA CGT TGCACTT GACTGC CGC TCGGAG CGAG GCCCCA CCAG AGAGAC CAGT GCACT CAGT GCACT CAGT GCACT CAGT GCACT CAGT GCACT CAGT GATTCA CCGC AATGGA AAG AAAGGAC TTG GATTAGG AAC CAAGGAC CAAC CCAGGGA CAACGAC CAACGGAC CAACGAC CCAC CCTGGG	CTGC 60 CTGGG 120 CGC 180 GCTAG 240 GCA 300 TGAGC 360 GTT 420 TGG 480 TGGT 540 GAGG 660 AGGG 660 TGCA 720 GATG 780 CCTIG 840 TGGGG 900 TTGTG 960 CCTC 1020 GGTT 1080 TTTA 1140 ACCT 1200 CACA 1260 GGCT 1320 TACA 1380	dons)
5055606570	SEQ ID NO: 188 E Nucleic Acid Acce 1 11 2 GCGGTCCGCC CCGCAACCGC CCGCAACCGC AGCCTGGCTA TCAATGGACT TCAATGGACT GCAGCGTCGGC TCTTGACCTG GCCCAGAGCG ATGACTGCCA AGCCTCGGCT AGAACAATGC GCAGCGTCGC TCTTGACCTG GCCCAGAGT ACCGCACCTC TCCAGTTGGA ATAACCGGC ACCGCACCTC TCCAGTTGGA ATTATTAAC GTGACCACAC CCTGTATGCG AGCGACGGC ACCTGCACCTC ACCGCACCTC ACCTCCACTCC ACCCCCCCCC ACCCCCCCCC	GCO2 DNA sequer ession #: Coding 1 31 4 G CACACCTCC C TGAGCCATC G CTGTGCCGC A TGCCGACGC A TGCCGACGC C CAAGGGGC C CATGTTGCC G CTGTTTCCT G CATGTTGCC G CTTTTTCCT G CATGAATAA C CTGTGAGTC G TAACCATGG G TAACCATGG G TAACCATGG G TAAGGATAC C TGCAAAAA C A CACTGTCC T GAAACGCC T GAAACGCC C TGCAAAAA C A TGGCAAGAC C TTCTCACCAC C TTCACCCA G CTTCACCCA G CTTCACCCA G CTTCACCCA G CTTCACCAA G CTTCACCCAA G CAATAAAA G CAAAAAA G CTGCATCAAA G CAAAAAA G CTGCATCAAA G CTGCATCAAA G CTGCATCAAA G CAAAAAAA G CTGCATCAAA G CTGCATCAA G CAAAAAAA C CAAGATGAAAAA C CAAGATGAAAAA C CAAAAAAA C CAAGATGAAAAA C CAAGATGAAAAAA C CAAGATGAAAAAA C CAAGATGAAAAA C CAAGATGAAAAA C CAAGATGAAAAAA C CAAGATGAAAAAA C CAAGATGAAAAA C CAAGATGAAAAAA C CAAGATGAAAAAA C CAAGATGAAAAAA C CAAGATGAAAAAA C CAAGATGAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAA	AJ400877 Sequence: 41 51 CC CGCGCCGC CC ATGGGGGT CC CTGCTGCCC CG GGGCCGCC CC CTGTGTCA AA GGCAGGCT T CATGACGGT AG CATACCT G AGTGACAA AG GATCACGC G AACGTGGA T CACACGT CT CACACGT CACACCT CACA	CG CCGCCAC CG CGGCCGC CG CGGCCGG CG AGGATGT GA ACACAC CG ATATTCCA TC ATAATTGT GT TCAACGT TTG TCAACGT TTG TCAACGT TTG TCAACGT TT TTGAGCAC TT TTGAGCAC TT TTGAGCAC TT TTGAGCAC TTG TCACTGC TCACTG TCACTGC TCACTG	1-3080 (underlined CGC CCGCACT CAA CCGTCCC CT GCTGGCG TAGA TGAGTG CACAT CGATGA CGT CAATTAT CT TGATGTGC CAT GGGGAG CTG CATTCAC CAT CTGCAAC GGC CAAGAAC CTC CTGTGAC CAC AGATGG TGC TGTCAAC CAG TTGTCCT GTG CCAGAC CTG CAGCAT CTGC CAGCTGC TTG CCAGCTG TTG CCAGCTC TTG TGCTTGC TTG CAGCATC CTG CAGCATC CTG CGCTGC TTG CAGCATC CTG CGCTGC TTG CAGCATC CTG CAGTGC CGG CTCCT	CCG CCGCCTC CGGG GCGCCC CGGG GCGCCC CGGG GCGCCC CGGG GCGCCC CGGG GCGCCC CGGG GCGCCC CGT GGAAAT GGAAT GGAGGC CGT TGCACT GGAGGC CGG TGGAGC CGG GCCCA CGAG GCTCG CGT GGATCA CAG GAGGG CTG GATCA CGAGGG CTG GATCA CCG CATGGA CAG CAACGA CCG CATGG CAAC CACGGG CCAC CCGGGG CCAC CCCGGGG CCAC AACGAAC CCAC CCCGGGG CCCC ACAGGG CCCC CCCGGGG CCCC ACAAGT	CTGC 60 CTGGG 120 CCGC 180 GCTAG 240 GCA 300 TGAGC 360 GTT 420 TGG 480 TGCT 540 GAGG 600 AGGG 660 TGCA 720 GATG 780 CCTTG 840 CCTG 840 CTGT 1020 GGTT 1080 CTTC 1020 GGTT 1080 CTTA 1140 ACCT 1200 ACA 1260 GGCT 1320 TACA 1380 GTGT 1440	dons)
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5055606570	SEQ ID NO:188 E Nucleic Acid Acce 1 11 2 GCGGTCCGCC CCGCAACCGC CGGTACTGCT CCGGTCGCT TCAATGGACT AGACAATGG GCAAGGAGG GCCTGAGCT AGACAATGGC TTTGACTG GCAAGGAGG ATAAACGG ATAAACGG ATAAACGG ATAAACGG ATAAACGG ACCGCACCTC TCCAGTTGGA GTGATCATTT AATTATAAC CCCTGTATGG GTCAGCAGGT CACCCCGTGT GTCACCCGGT GTCACCCGGG GTCACCCGGG GTCACCCGGGG GTCACCCCGTGT GTCACCCGGG GTCACCCGGGG GTCACCCCGTGT GTCACCCCGTGT	BCO2 DNA sequer Ession #: Coding 21 31 4 G CACACCTCC C TGAGCCATC G CTGCTGCTG G CCGTGCTGCT G CCGTGCTGCT G CATGTTGCC G GATTTTCCT G CATGATGC G TAACCATGG G TAACCATGG G TAACCATGG G TAACCATGG G TAACCATGG G TAAGGAAA C TGGAAAAA C TGGAAAAA C TGGCATCACCA G CTTCACCA G CTTCACCAC G CATTCACCAC C C CATTCACCAC C C C C C C C C C C C C C C C C C	AJ400877 Sequence: 41 51 CC CGCGCCGC CC ATGGGGGT GC TGCTGCCC CC GGGCCCGC CC CTGTGTCA AA GGCAGGC AT GACTGTTCA AG CATACCTC GG AGTGACAA AG GATACCTC GG AGCCTGGG AT CCACAGTA TT CCACAGTA TT TCGACAGG CA TGTAAAGA AC ATCGTGGG AG TCTTGCCA AC CACCCTGG CC TGTGGAGAGA AC CACCTTGG CC TGTGGAGAGA AC CACCTTGC CC TGTGGAGAGA AC CACCTTGC CC TGCGGTAAA	CG CCGCCAC CG CGGCCAC CG CGGGCCG AGG AGGATGT GA ACACACC AGT GTGAGG AGCACAC TC ATAATTGT TG TCAACGT TT TTGAGCAC AGT GTAGCAC AGT GCAGCAC AGT GCACCAC AGT GCACCAC AGT TTGATGAC AG ATGTGGA AG AGTGGA AG AGTGGA AG AGTAGAC AG AGTAGAC AG AGTGTGAC AG AGTGTGAC AGG AGTGGAC AGG AGTGGAC AGG AGTGGGAC AGG AGTGGGAC AGG TCACCACC	1-3080 (underlined CGC CCGCACT CAA CCGTCCC CTG CTGCGGG CAGA TGAGTG CAC CTCCTAC ACAT CGATGA CGC CAATTAC CAT GGGGAG CTG CATTCAC CAT CTGCAAA CTC CTGTGAC CAC AGATG CAC AGATG CAC AGATG CAC GGGCTGC CTG CGGCTGC TGG TGCTCT TGG TGCTCT TGG TGCTCT TGG CAGGAC ATG CCAGTGG ATG CCAGTGG ATG CCAGTGG ATG CCAGTGC ATG CAGGAC ATG CAGGAC ATG CAGGAC ATG CAGGAC ATG CAGGAC ATCAGGACA ATCAGGACA ATCAGGACA	CCCG CCGCCTC CCGG GCGCCTC CCGG GCCGCCTC CCGG GCCGCCC CCGG GCCGCCC CCAAGG CAAG TGCTCCT CAAGT GCAAA CCGA GCGCC CCAT GAGTGCC CCAT GAGTGC CCAT GAGTGC CCAT GAGTGC CCAT GAGTGC CAGT GCACT CAAT GAAGAC CCAT GAATGA CCAT GAATGA CCAT GAATGA CCAT GAATGA CCAT GAATGA CCAC CACAGG CCCC ACAGG CCCC ACAGG CCCC ACAGG CCCC ACAGG CCCC ACAGG CCCC ACAAGT CCCC CCAAGT CCCC CCACAGT CCCC CCACAGT CCCC CCCACAGT CCCC CCCCC CCCCC CCCCC CCCCC CCCCC CCCCC CCCC	CTGC 60 CTGGG 120 CGC 180 GCTAG 240 GCA 300 TGAGC 360 GTT 420 TGG 480 TGGC 540 GAGG 660 TGCA 720 GATG 780 CCTTG 840 CCTTG 840 CTGTG 960 CTC 1020 GGTT 1080 TTTA 1140 ACCT 1200 CACA 1260 GGCT 1320 TACA 1380 GGTGT 1440 AGAT 1500 TTA 1500	dons)
5055606570	SEQID NO: 188 E Nucleic Acid Acce 1 11 2 GGCGTCCGCC CCGCAACCGC CCGCAACCGC CGGTCGGGCT AGACCAGCT AGACAATGGCT AGACAATGGCT AGACAATGGCT GCAAGCAGG GCCAGAGT GCAGAGT AGCGAGAGG ATAAACGGG ATAAACGGG ATAATTATTAAC GTGACCACAC CCCTGTATGC GTCAGCAGGG AGCTCCACTG AGCTAAATGA	GCO2 DNA sequer sesion #: Coding 21 31 4 G CACACCTCC G GCACACCTCC G CCGTGCCGC A TGCCGACGC G CATGTTGCCA G CATGTTGCCA G CATGTTGCC G CATGTTGCC G CATGTTGCC G CATGTTGCC G CATGTTGCC G CATGTAAAAA C CTGTGAGTA C CATGTCCC T GAAACAGC G CATGCCCA A TGCAAAAA C AGATGAGAA C CTGCATCAA C CTGTGTGCAA C CTGTGTGCAA C CTGTGTCACCAA C CTGTGTCACCAA C CTGTGTCACCAA C CTGTGTCACCAA C CTGTGTCACCAA C CTGTGTCACCAA C CATTCACCCT C CATTCACCCT C C CATTCACCCT C C C C C C C C C C C C C C C C C C	AJ400877 Sequence: 41 51 CC GCGCCGC CC ATGGGGGT GC CTGTGTCA AA GGCAGGC AT GACTGTTT T CATGACGGT AG CATACCT GG GATCACAG AG GATCACAG AG GATCACAG AT CACAGTA AT GACGTGA AG CATCACAG AT CACAGTA AG CATCACAG AT CACAGTA AG CATCACAG AT CACAGTA AG CATCACAG AT CTACAGG AG CTGTCAA AC CACCTGG AG TCTTGCA AC CACCTGG AG TCTTGCAA AC CACCTGG AG TCTTCAAA AC CACCTGG AC TCTTCAAA AG CTCTCAAA AG CTCTTCAAA AG TCTTCAAA AG TCTTCAAA AG TAGTTTGAA	CG CCGCCAC CCG CCGCCAC CCG CCGCCCAC CCG CCGCCCC CGC CCGCCCC CGC CCGCCCC CGC CGCCCC CGC CGCCCC CGC CGCCCC CGC CCCCCC CCGCCCCC CCGCCCCCCCC	1-3080 (underlined CGC CCGCACT CAA CCGTCCC CAG TGGGGG TAGA TGAGTG CAC TCCTAC ACAT CGATGA CGC CATTAT CT TGATGTG CAT CGGGAG CTG CATTCAC CAT CGGGAG CTG CATTCAC CAT CTGCAAC CAC TGTGAAC CAC TGTGAAC TGC TGTCAAC TGC TGCAAC TGC TGCAAC TGC CAGAC TGC CAGAC TGC CAGAC TG CCGCTGC TG CGGCTGC TG CGGCTCC TG CGGCTCC TG CGGCTCC TG CGGCTCC TG CAGCAC CAGAC CAGGAC CAGC CAGGAC CAGC CAGGAC CAGGAC CAGGAC CAGGAC CAGGAC CAGGAC CAGGAC CAGGAC CAGC CAGAC CAGGAC CAGC CAGGAC CAGC CAGC CAGGAC CAGC CAGGAC CAGC CAC CA	CCG CCGCCTC CGG CCGCCTC CGGG GCGGCC GGG GCGGCC CTGC CAAGGC CAG TGCACTT GACTGCAC CGC TGCACTC CGC TGCACTC CGC TGCACTC CGC TGCACT CGC AGGG CGC ACAGG CGC AATGGA CAG CGACGA CAG CGACGA CAG CGACGA CACCC CAGT GATACA CCGC CATGGA CCCC CACGG CCAC CCTGGG CCCC CACAGT CCAC CCTGGG CCCC ACAGT CGGG GTTCACC CGGG GCCC ACAGT CGGG GTTAACCT CGAG GGTCTGC	CTGC 60 CTGGG 120 CGC 180 GCTAG 240 GCA 300 TGAGC 360 GTT 420 TGG 480 TGGC 540 GAGG 660 TGGA 720 GATG 780 CCTIG 840 CCTIG 960 CCTC 1020 GGTT 1080 TTTA 1140 ACCT 1200 TACA 1260 GGCT 1320 TACA 1380 GTGT 1440 AGGT 1500 TTA 1500 CTA	dons)
5055606570	SEQID NO: 188 E Nucleic Acid Acce 1 11 2 GCGGTCCGCC CCGCAACCGC CCGCAACCGC AGCCTGGCT AGACATGC AGCAGAGAGG GCTGAGCT GCAAGCAGG TTGATGCTT AGAACAATG GCAAGCAGG GCCTAGACT GCCAGATG GCCAGAGT AGCACTCG ATTATTAAC GTGACCACT CCAGTTGGC AGTCACT CCAGTTGGC AGCACACT CCAGTTGAC CCCTGTATTG ATTATTAAC CCCTGTATTG ACCCCGTGT CCACCCCGTGT CCACCCCCGTGT CCACCCCCGTGT CCACCCCCGTGT CCACCCCCCCCCC	GCO2 DNA sequer ession #: Coding 21 31 4 G CACACCTCC C TGAGCCATC G CTGTGCCGC A TGCCGACGC A TGCCGACGC G CTGTTTTTCC G CTGTGAGTCC G CATGAGTCC G CATGAGTA C CTGCAAAAA C CTGCAAAAA C ACACTGCC G TGAGCAC G CTTCACCAC G CATTCACCC G AGGAAAAA C GTCCCTGCA G CATTCACCC G AGGCAAGTC C AGGCAAGTC C AGGCAAGTC C CATCCCC C CACCTGCC C CACCTGCC C CACCTGCC C CACCTGCC C CACCTGCC C CACCTGC C C CACCTGC C C C C C C C C C C C C C C C C C C C	AJ400877 Sequence: 41 51 CC CGCGCCGC CC ATGGGGGC CC GGGCCGCC CC GGGCCGCC CC GGGCCGCC CC GGGCCGCC CC GGGCCGCC CC GGGCCGCC CC CTGTGTCA AA GGCAGGCT AG CATACCTC G AGGCAGCA AG GATCACGC GG AACGGTGG AC ATGCACAGC CC ATGGACAA CC ATCGTGGGC CC TGTGGAGA AC GCTCTCCA AC CACCTGGG CC TGTGGAGA AC CACCTGGG CC TGTGGAGA AC CACCTGGG AC GCTCTCAC AC CACCTGGG AC GCTCTCAC AC CACCTGGG AC TCTTCACAG AC AGTTGGAA AC AGCTCAGT AGTTTTCAAA AC AGCTCAGT AC	CG CCGCCAC CG CGGCCAC CG CGGCCGG CG CGGCCGG CG AGGATGT GA ACACAC CG ATATTCCA TC ATAATTGT GT CAACACC CT GTGAGGA TT TTGAGCT GT GCCAGCA CT GTGAGCA CA AGATGCA CA AGATGCA CA CACTG AGA ACC TG TCACTG CA CACTG CA CATTTGC CA CAATTGC CAAATGC CAAAAAGAGAG CAAAAAGAGAG	1-3080 (underlined CGC CCGCACT CAA CCGTCCC CAA CCGTCCC CAC TGATGGCG CAC TCGATGA CACAT CGATGG CAT GATGAC CAT GATGAC CAT GATGAC CAT CGATGAC CAC CACTCAC CAC CACTCAC CAC CACTCAC CAC CACTCAC CAC TGCAAC CAC TGCAAC CAC TGCAAC CAC TGCAAC CAC TGCAAC CAC TGCCAAC CAC TGCCAC CTG CAGCTC CTG CCACTC CTG CAGCTC CTG CAGCTC CTG CAGCTC CTG CAGCTC CTG CAGCTC CACTC CACTC CCTA	CCG CCGCCTC CCGG CCGCCTC CCGG GCGGCCC CTGC CAAGGC CAAG TGCTCCT CGT TGCACTT GAC GAGTGCC CTAT GAGTGCC CTAT GAGTGCC CTAT GAGTGCC CTAT GAGTGCC CTAT GAGTGCC CTAT GAGTGCC CAGT GCACC CAGT GCACC CAGT GCACC CAGT GATCA CCGC AATGGA CCGC AATGGA CCGC AATGGA CCGC CATGGA CAC CCTGGG CCCC CCGGG CCCC CCGGG CCCC CCGGG CCCC CCGGG CCCC CCAGGGT CAGC GTTCCTCC CGAG GGTCTG CCGTA AACCTT	CTGC 60 CTGGG 120 CTGGG 120 CCGC 180 GCTAG 240 GCA 300 TGAGC 360 GTT 420 TGG 480 TGGT 540 GAGG 660 AGGG 660 TGCA 720 GATG 780 CCTTG 840 CCTG 840 CTGT 1020 GGTT 1080 CTTC 1020 GGTT 1080 CTTA 1140 ACCT 1200 ACA 1260 GGCT 1320 TACA 1380 GTGT 1440 AGAT 1500 CTGAC 1620 ACAT 1680	dons)
505560657075	SEQIDNO:188 E Nucleic Acid Acce 1 11 2 GCGGTCCGCC CCGCAACCGC CCGCAACCGC CGGGTCCGGCT CGGGTCGGCT TCAATGGAG TTGATGGCT AGAACAATGG GCAAGGAGGG GCTGAGCTC GCCCAGAGTC GCAGCGTCGG TCTTGACCTG GCCAGAGTC AGCAGAGGG ATTATTAAC GTGACCACCC CCTGTATGG GTCAGCAGG GTCAGCAGG GTCAGCAGG GTCAGCAGG GTCAGCAGG GTCACTCG CACCCGTGT GTCACTCG CACCCGTGT GTCACTCG CACCCGTGT GTCACTCG CACCCGTGT GTCACTCG CACCCGTGT GTCACTCG CACCCGTGT GTCACTCG CAGCACTACC CAGCACTCC	GCO2 DNA sequer ession #: Coding 1 31 4 G CACACCTCC C TGAGCCATC G CTGCTGCTGCT G CCGTGCCGC A TGCCGACGC A TGCCGACGC C CAAGGGGA C CAAGGGGA G CATGTTGGC G CGGTTGCCA T CATGTTGGC G TTTTTCCT T GAAACGGC T TAACCATGG T TAACCATGG T CATGATAC C TGGAAACG T CTGCAAAA C ATGCAAAA C ATTCACCCA C AGATGACCA C AGATAAAAA C GTCCCTGCA C AGAGCAAGT C AGAGCAAGT C AGAGCAAGT C AGAGCAAGT C AGAGCAAGT C AGAGCAAGT C CAAGCAAGT C C CAAGCAAGT C C CAAGCAAGT C C C C C C C C C C C C C C C C C C C	AJ400877 Sequence: 41 51 CC CGCGCCGC CC ATGGGGGT CC ATGGGGGT CC CTGTGTCA AA GGCAGGC AG CATACCT CG AGGCCTGG AG CATACCT CG AGGCCAGG AG CATACCT CG AGGCTGA AT CCACAGTA AT GACGGTGA AG CATCGTGG CA TGCTCAT CT TCGACAGG CA TGCTCAT CT TCGACAGG AG CTTGCCA CC ATCGTGGA AC CACCTGG CC TGTGGAGA AC CACCTGG CC TGTGGAGA AC CACTTGGT CT TCGAGAGA AC AGCTGAGT CT TCGAGAGA AC AGCTCAGT CT TCCAGGAGA AC AGCTCAGT CC CCAGGAGA CC CCAGGAGAG CC CCCCAGGAGAG CC CCCCAGGAG CC CCCCCAGGAG CC CCCCCAGGAG CC CCCCCAGGAG CC CCCCCAGGAG CC CCCCCAGGAG CC CCCCCAGGAG CC CCCCCCAG CC CCCCCCC CCCCCCCCCC	CG CCGCCAC CG CGGCCAC CG CGGCCGG CG CGGGCCGG CG AGGATGT GA ACACACC AGT GTGAGG, GA ATATTCCA TC ATAATTGT GTG TCAACGT TTGAGCAC CT GTAGTCA CT TTGAGCAC CA AGATGCA AGT TTGAGCAC CA GATGTGA CA CATTTGA CA CA	1-3080 (underlined CGC CCGCACT CAA CCGTCCC CT GCTGGCG CAG TGAGTG CACAT CGATGA CGT CAATTAT CT TGATGTGC CAT GGGGAG CTG CAATTAAC CAT CGGAGA CAC CACAT CGC CAAGAAC CTC CTGTGAC CAC GGC CAAGAAC CTC CTGTGAC CAC GGCTGC TGC CAGACT CTG CTC CTG CTC CTG CTC	CCCG CCGCCTC CCGG GCGCCTC CCGG GCCGCCTC CCGG GCCGCCC CTGCC CAAGGC CAAG TGCTCCT CGT TGCACTT CGACAT CAG AGGGCC CTAT GAGTGC CCGC TCGGAAG CCAG AGGGCC CAGG GCCCCA CCAG AGGGCC CAT GGAGT CAGT GCGCC CAGT GCGCC CAGT GCGCC CAGT GCGCC CAGT GCGCC CCGC CATGGA CCC CCTGGG CCC CACAGT CCCC CACAGT CCCC CACAGT CCCC CACAGT CCCC ACAGT CCCC ACAGT CCCCT CCCCC CCGCC CCGTA ACCCTT CCCT AAGGAA	CTGC 60 CTGGG 120 CGC 180 GCTAG 240 GCA 300 TGAGC 360 GTT 420 TGG 480 TGGT 540 GAGG 600 AGGG 660 TGGA 720 GATG 780 CCTTG 840 TGGAT 780 CCTG 1020 GGTT 1080 TTATA 1140 ACAT 1200 TACA 1260 GGGT 1320 TACA 1380 GTGT 1440 AGAT 1500 TTA 1500 CCAC 1620 ACAT 1680 AATGT 1740	dons)
5055606570	SEQIDNO:188 E Nucleic Acid Acce 1 11 2 GCGGTCCGCC CCGCAACCGC CCGCAACCGC CGGGTCCGGCT CGGGTCGGCT TCAATGGAG TTGATGGCT AGAACAATGG GCAAGGAGGG GCTGAGCTC GCCCAGAGTC GCAGCGTCGG TCTTGACCTG GCCAGAGTC AGCAGAGGG ATTATTAAC GTGACCACCC CCTGTATGG GTCAGCAGG GTCAGCAGG GTCAGCAGG GTCAGCAGG GTCAGCAGG GTCACTCG CACCCGTGT GTCACTCG CACCCGTGT GTCACTCG CACCCGTGT GTCACTCG CACCCGTGT GTCACTCG CACCCGTGT GTCACTCG CACCCGTGT GTCACTCG CAGCACTACC CAGCACTCC	GCO2 DNA sequer ession #: Coding 1 31 4 G CACACCTCC C TGAGCCATC G CTGCTGCTGCT G CCGTGCCGC A TGCCGACGC A TGCCGACGC C CAAGGGGA C CAAGGGGA G CATGTTGGC G CGGTTGCCA T CATGTTGGC G TTTTTCCT T GAAACGGC T TAACCATGG T TAACCATGG T CATGATAC C TGGAAACG T CTGCAAAA C ATGCAAAA C ATTCACCCA C AGATGACCA C AGATAAAAA C GTCCCTGCA C AGAGCAAGT C AGAGCAAGT C AGAGCAAGT C AGAGCAAGT C AGAGCAAGT C AGAGCAAGT C CAAGCAAGT C C CAAGCAAGT C C CAAGCAAGT C C C C C C C C C C C C C C C C C C C	AJ400877 Sequence: 41 51 CC CGCGCCGC CC ATGGGGGT CC ATGGGGGT CC CTGTGTCA AA GGCAGGC AG CATACCT CG AGGCCTGG AG CATACCT CG AGGCCAGG AG CATACCT CG AGGCTGA AT CCACAGTA AT GACGGTGA AG CATCGTGG CA TGCTCAT CT TCGACAGG CA TGCTCAT CT TCGACAGG AG CTTGCCA CC ATCGTGGA AC CACCTGG CC TGTGGAGA AC CACCTGG CC TGTGGAGA AC CACTTGGT CT TCGAGAGA AC AGCTGAGT CT TCGAGAGA AC AGCTCAGT CT TCCAGGAGA AC AGCTCAGT CC CCAGGAGA CC CCAGGAGAG CC CCCCAGGAGAG CC CCCCAGGAG CC CCCCCAGGAG CC CCCCCAGGAG CC CCCCCAGGAG CC CCCCCAGGAG CC CCCCCAGGAG CC CCCCCAGGAG CC CCCCCCAG CC CCCCCCC CCCCCCCCCC	CG CCGCCAC CG CGGCCAC CG CGGCCGG CG CGGGCCGG CG AGGATGT GA ACACACC AGT GTGAGG, GA ATATTCCA TC ATAATTGT GTG TCAACGT TTGAGCAC CT GTAGTCA CT TTGAGCAC CA AGATGCA AGT TTGAGCAC CA GATGTGA CA CATTTGA CA CA	1-3080 (underlined CGC CCGCACT CAA CCGTCCC CT GCTGGCG CAG TGAGTG CACAT CGATGA CGT CAATTAT CT TGATGTGC CAT GGGGAG CTG CAATTAAC CAT CGGAGA CAC CACAT CGC CAAGAAC CTC CTGTGAC CAC GGC CAAGAAC CTC CTGTGAC CAC GGCTGC TGC CAGACT CTG CTC CTG CTC CTG CTC	CCG CCGCCTC CCGG GCGCCTC CCGG GCCGCCTC CCGG GCCGCCC CTGC CAAGGG CAAG TGCTCCT AAGT GGAAA CGAT TGCACTT GAC GAGTGCC CTAT GAGTGC CCTAT GAGTGC CCGAGG GCCCCAAC CCAG GGAGAC CAAT GGAGG CCCTAT GAGTGC CAAT GGAGG CCGAAC CAAT GGAGG CCTAT GAGTGC CAAT GGAGG CCTAT GAATCA CCGAAT GCACC CGGG ACAGGG CTTG GATACA CCGAGGGT CAAC CCTGGG CCCC ACAAGT CCCCT AAGGAA CCCTTCCCCT CCCT AAGGAA CCCT TCCTTCCCCT CCCT AAGGAA CCCT TCCTTCTCCCCT CCCT AAGGAA CCCT TCCTTCTCCCT CCCT AAGGAA CCCT TCCTTCTCTCCCT CCCT TCCTTCTCTCTC	CTGC 60 CTGGG 120 CGC 180 GCTAG 240 GCA 300 TGAGC 360 GTT 420 TGG 480 TGGT 540 GAGG 600 AGGG 660 TGGA 720 GATG 780 CCTTG 840 TGGAT 780 CCTG 1020 GGTT 1080 TTATA 1140 ACAT 1200 TACA 1260 GGGT 1320 TACA 1380 GTGT 1440 AGAT 1500 TTA 1500 CCAC 1620 ACAT 1680 AATGT 1740	dons)

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GTTCACCTGG ACATTTCTAC AACACCACCA CTCACCGATG TATTCCTTGC CCAGTGGGAA 2400 10 CATACCAGCC TGAATTTGGA AAAAATAATT GTGTTTCTTG CCCAGGAAAT ACTACGACTG 2460 ACTITGATGG CTCCACAAAC ATAACCCAGT GTAAAAACAG AAGATGTGGA GGGGAGCTGG GAGATTTCAC TGGGTACATT GAATCCCCAA ACTACCCAGG CAATTACCCA GCCAACACCG 2580 AGTGTACGTG GACCATCAAC CCACCCCCA AGCGCCGCAT CCTGATCGTG GTCCCTGAGA 2640 15 TCTTCCTGCC CATAGAGGAC GACTGTGGGG ACTATCTGGT GATGCGGAAA ACCTCTTCAT 2700 CCAATTCTGT GACAACATAT GAAACCTGCC AGACCTACGA ACGCCCCATC GCCTTCACCT 2760 CCAGGTCAAA GAAGCTGTGG ATTCAGTTCA AGTCCAATGA AGGGAACAGC GCTAGAGGGT 2820 TCCAGGTCCC ATACGTGACA TATGATGAGG ACTACCAGGA ACTCATTGAA GACATAGTTC 2880 TCAGGICCE ATACGIGACA TATGATGAGG ACTACCAGGA ACTCATIGAA GACATAGTTC 2880
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GAACTTGGTT TTTCTTTCCC AGCATCGTGG ATGTAGACTG AGAATGGCTT TGAGTGGCAT 3300 20 25 CAGCTTCTCA CTGCTGTGGG CGGATGTCTT GGATAGATCA CGGGCTGGCT GAGCTGGACT 3360 TTGGTCAGCC TAGGTGAGAC TCACCTGTCC TTCTGGGGTC TTACTCCTCC TCAAGGAGTC 3420 TGTAGTGGAA AGGAGGCCAC AGAATAAGCT GCTTATTCTG AAACTTCAGC TTCCTCTAGC 3480 CCGGCCCTCT CTAAGGGAGC CCTCTGCACT CGTGTGCAGG CTCTGACCAG GCAGAACAGG 3540 30 CAAGAGGGGA GGGAAGGAGA CCCCTGCAGG CTCCCTCCAC CCACCTTGAG ACCTGGGAGG 3600 ACTCAGTTTC TCCACAGCCT TCTCCAGCCT GTGTGATACA AGTTTGATCC CAGGAACTTG 3660 AGTTCTAAGC AGTGCTCGTG AAAAAAAAA GCAGAAAGAA TTAGAAATAA ATAAAAACTA 3720 AGCACTTCTG GAGACAT 35 SEQ ID NO:189 BCO2 Protein sequence CAB92285 Protein Accession #: 21 31 40 45 STGVHCSCPV GFTLQLDGKT CKDIDECQTR NGGCDHFCKN IVGSFDCGCK KGFKLLTDEK 360 SCODVDECSL DRTCDHSCIN HPGTFACACN RGYTLYGFTH CGDTNECSIN NGGCQQVCVN 420 TVGSYECQCH PGYKLHWNKK DCVEVKGLLP TSVSPRVSLH CGKSGGGDGC FLRCHSGIHL 480 SSDVTTIRTS VTFKLNEGKC SLKNAELFPE GLRPALPEKH SSVKESFRYV NLTCSSGKQV 540 50 PGAPGRPSTP KEMFITVEFE LETNQKEVTA SCDLSCIVKR TEKRLRKAIR TLRKAVHREQ 600 FHIQLSGMNL DVAKKPPRTS ERQAESCGVG QGHAENQCVS CRAGTYYDGA RERCILCPNG 660
TRONEEGQMT CEPCPRPGNS GALKTPEAWN MSECGGLCOP GEYSADGFAP CQLCALGTFQ 720
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PPPKRRILIV VPEIFLPIED DCGDYLVMRK TSSSNSVTTY ETCQTYERPI AFTSRSKKLW 900 55 IQFKSNEGNS ARGPQVPYVT YDEDYQELIE DIVRDGRLYA SENHQEILKD KKLIKALFDV 960 LAHPONYFKY TAQESREMFP RSFIRLLRSK VSRFLRPYK SEQ ID NO:190 BFG1 DNA sequence 60 AF007170 Nucleic Acid Accession #: Coding sequence: 1-1725 (underlined sequences correspond to stop codon) 41 31 AAGGAGGCGG CCTCCGGGAA AAGCGACCGC AGGACTCCTG AGAGCAGCCT CCATGAGGCC 60
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ATGAAGGAGG CACAGATGCT GTGTCAGAGG CACCGGAGGA AGTCTTCTGT AACAGATTCC 300 65 70 TTCAGCAGCC TGGTGAACCG CCCCACGCTG GGCCAATTCA CTGAAGAAGA AATCCACGCT GAGGTCTGCT ATGCAGAGTG CCTGCTGCAG CGAGCAGCCC TGACCTTCCT GCAGGACGAG 420 AACATGGTGA GCTTCATCAA AGGCGGCATC AAAGTTCGAA ACAGCTACCA GACCTACAAG 480
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CTGTGCTACC ACACCTTCCT CACCTTCGTG CTCGGTACTG GGAACGTCAA CATCGAGGAG 780 75 GCCGAGAAGC TCTTGAAGCC CTACCTGAAC CGGTACCCTA AGGGTGCCAT CTTCCTGTTC 840
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GAGTGCTGTG AGGCCCAGCA GCACTGGAAG CAGTTTCACC ACATGTGCTA CTGGGAGCTG 960 ATGTGGTGCT TCACCTACAA GGGCCAGTGG AAGATGTCCT ACTTCTACGC CGACCTGCTC 1020 AGCAAGGAGA ACTGCTGGTC CAAGGCCACC TACATTTACA TGAAGGCCGC CTACCTCAGC 1080 ATGITTGGGA AGGAGGACCA CAAGCCGTTC GGGGACGACG AAGTGGAATT ATTTCGAGCT 1140
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CGGAAGTCCC GGCGCTACTT CTCCTCCAAC CCTATCTCGC TGCCAGTGCC TGCTCTGGAA 1260 5 ATGATGTACA TCTGGAACGG CTACGCCGTG ATTGGGAAGC AGCCGAAACT CACGGATGGG 1320 ATACTTGAGA TTATCACTAA GGCTGAAGAG ATGCTGGAGA AAGGCCCAGA GAACGAGTAC 1380 TCAGTGGATG ACGAGTGCTT GGTGAAATTG TTGAAAGGCC TGTGTCTGAA ATACCTGGGC 1440 10 CGTGTCCAGG AGGCCGAGGA GAATITTAGG AGCATCTCTG CCAATGAAAA GAAGATTAAA 1500 TATGACCACT ACTTGATCCC AAACGCCCTG CTGGAGCTGG CCCTGCTGCT TATGGAGCAA 1560 GACAGAAACG AAGAGGCCAT CAAACTTTTG GAATCTGCCA AGCAAAACTA CAAGAATTAC 1620 TOCATGGAGT CAAGGACACA CTTTCGAATC CAGGCAGCCA CACTCCAAGC CAAGTCTTCC 1680 CTAGAGAACA GCAGCAGATC CATGGTCTCA TCAGTGTCCT TGTAGCTTTG TGCAGCAGTT 1740
CCGGGCTGGA AGACAGAGAC AGCTGGACAG AGCTCCTGAA AACATTTCAA AATACCCCCT 1800
CCCCTGCCC TGCCCTGCCT TTGGGGTCCA CCGGCACTCC AGTTGGATGG CACAACATAG 1860
TGTATCCGTG CAGAAGCCGA GCTGGCATTT TCACCAGTGT AGCCAAGGGC CTTTGCCAAG 1920
GGCAGAGCAG GTGGAGCCCT CTGCCTGCCC TATCACACAT ACGGGTACTT GCTTTTCACT 1980 15 GTGATGTTTA AGAGAATGTA TGAACAGTTT ACATTTTCCT TAGAAATACA TTGATGGGAT 2040 20 CACAGITGGC TITAAAAACC AACAACAATC AACCACCTGT AAGTCTTTGT CTTCACCTAT 2100 TATCATCTGG AGGTAAATCT CTTTATATGA TGATGCCAAA GGGCAAATTG CTTTTCAAAT 2160 TCAGCAAGTT CTCAGCTTGT GTGACGGAAG GTCCTTCAGA GGACCTGAGG AATGCCTGGG 2220 AGAGGCTAAG CCTCAGGCTT CAATGCTTCT GGGGTTGGGC ATGAGGATGT ACACAGACAC 2280 CCACTACCTT ACTACTCACA CTTCATTTCA CTCCTTTTGT AAATTTCCAA TTTAAAAATC 2340 25 AAGCACGTCT TTTTAGTGAG ATAAAATCTG AGCTCTTCTG TAGAAAAATC AATCTCTACC 2400 AGTAGAAAAT GCCAGGGCTT GATGGAAGAG CTGTGTAGCC CTTTCTATGC CAAAGCCAGG 2460 AAATTTGGGG GGCAGGAGGA GGTTCTCAGA ATCCAGTCTG TATCTTTGCT GTATGCCAAA 2520 CTGAAACCAC TGGGAATAAT TTATGAAACA TAAAAATCTT CTGTACTTCA CTCCAAGGTA 2580 CATTIGCTTA CIGACAGCAT TITIGTTAAA ACTGTTATTC TIGAAAAAAA AAAAAAAAA 2640 30 SEQ ID NO:191 BFG1 Protein sequence AAC39582 Protein Accession #: 35 41 MTALDLFITN QFSEALSYLK PRTKESMYHS LTYATILEMQ AMMTFDPQDI LLAGNMMKEA 60
QMLQRHRRK SSVTDSFSSL VNRPTLGQFT EEEHAEVCY AECLLQRAAL TFLQDENMVS 120
FIKGGIKVRN SYQTYKELDS LVQSSQYCKG ENHPHFEGGV KLGVGAFNLT LSMLPTRILR 180
LLEFVGFSGN KDYGLLQLEE GASGHSFRSV LCVMLLLCYH TFLTFVLGTG NVNIEEAEKL 240
LKPYLNRYPK GAIFLFFAGR IEVIKGNIDA AIRRFEECCE AQQHWKQFHH MCYWELMWCF 300 40 TYKGQWKMSY FYADLLSKEN CWSKATYIYM KAAYLSMFGK EDHKPFGDDE VELFRAVPGL 360 KLKIAGKSLP TEKFAIRKSR RYFSSNPISL PVPALEMMYI WNGYAVIGKQ PKLTDGILEI 420 45 ITKAEEMLEK GPENEYSVDD ECLVKLLKGL CLKYLGRVQE AEENFRSISA NEKKIKYDHY 480 LIPNALLELA LILMEQDRNE EAIKLLESAK QNYKNYSMES RTHFRIQAAT LQAKSSLENS 540 SRSMVSSVSL 50 SEQ ID NO:192 BFO6 DNA sequence Nucleic Acid Accession #: NM 032583 Coding sequence: 1-4044 (underlined sequences correspond to start and stop codons) 21 31 41 55 ATGACTAGGA AGAGGACATA CTGGGTGCCC AACTCTTCTG GTGGCCTCGT GAATCGTGGC 60 ATCGACATAG GCGATGACAT GGTTTCAGGA CTTATTTATA AAACCTATAC TCTCCAAGAT 120 GGCCCCTGGA GTCAGCAAGA GAGAAATCCT GAGGCTCCAG GGAGGGCAGC TGTCCCACCG 180 TGGGGGAAGT ATGATGCTGC CTTGAGAACC ATGATTCCCT TCCGTCCCAA GCCGAGGTTT 240 60 CCTGCCCCC AGCCCCTGGA CAATGCTGGC CTGTTCTCCT ACCTCACCGT GTCATGGCTC 300
ACCCCGCTCA TGATCCAAAG CTTACGGAGT CGCTTAGATG AGAACACCAT CCCTCCACTG 360
TCAGTCCATG ATGCCTCAGA CAAAAATGTC CAAAGGCTTC ACCGCCTTTG GGAAGAAGAA 420
GTCTCAAGGC GAGGGATTGA AAAAGCTTCA GTGCTTCTGG TGATGCTGAG GTTCCAGAGA 480
ACAAGGTTGA TTTTCGATGC ACTTCTGGGC ATCTGCTTCT GCATTGCCAG TGTACTCGGG 540 65 CCAATATIGA TTATACCAAA GATCCTGGAA TATTCAGAAG AGCAGTTGGG GAATGTTGTC 600 CATGGAGTGG GACTCTGCTT TGCCCTTTTT CTCTCCGAAT GTGTGAAGTC TCTGAGTTTC 660 TCCTCCAGTT GGATCATCAA CCAACGCACA GCCATCAGGT TCCGAGCAGC TGTTTCCTCC TTTGCCTTTG AGAAGCTCAT CCAATTTAAG TCTGTAATAC ACATCACCTC AGGAGAGGCC 780 ATCAGCTTCT TCACCGGTGA TGTAAACTAC CTGTTTGAAG GGGTGTGCTA TGGACCCCTA 840 70 GTACTGATCA CCTGCGCATC GCTGGTCATC TGCAGCATTT CTTCCTACTT CATTATTGGA 900 GTACTGATCA CCIGCGCATC GETGGTCATC TGCAGCATTT CTTCTACTTACTT CATTATTGGA 900
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GAGCTGGAGA GGAACGGGCA TGCTTCTGAG GGGATGACCA GGCCTAGAGA TGCCCTCGGG 1440



WGKYDAALRT MIPFRPKPRF PAPQPLDNAG LFSYLTVSWL TPLMIQSLRS RLDENTIPPL 120 WGKYDAAIRT MIPFRYKRF PAPQPLDNAG LFSYLITVSWL TPLMIQSIRS RIDENTIPPL 120
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PILJIPKILE YSEEQLGNVV HGVGLCFALF LSECVKSLSF SSSWIINQRT AIRFRAAVSS ,240
FAFEKLIQFK SVIHITSGEA ISFFTGDVNY LFEGVCYGPL VLITCASLVI CSISSYFIG 300
YTAFIAILCY LLVFPLAVFM TRMAVKAQHH TSEVSDQRIR VTSEVLTCIK LIKMYTWEKP 360
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LQESPVFVQ TLQDFSKALV FEEATLSWQQ TCPGIVNGAL ELERNGHASE GMTRPRDALG 480
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YVPQQAWIVS GNIRENILMG GAYDKARYLQ VLHCCSLNRD LEILPFGDMT EIGERGLNLS 600
GGGVGDISLA PAVYSDROJV IL DDPI SAVD AHVGKHIEGE CIKKTI BGKT VM VYTRIOLOG 660 60 65 GGQKQRISLA RAVYSDRQIY LLDDPLSAVD AHVGKHIFEE CIKKTLRGKT VVLVTHQLQY 660 LEFCGQIILL ENGKICENGT HSELMQKKGK YAQLIQKMHK EATSDMLQDT AKIAEKPKVE 720 SQALATSLEE SLNGNAVPEH QLTQEEEMEE GSLSWRVYHH YIQAAGGYMV SCIIFFFVVL 780 70 IVFLTIFSFW WLSYWLEQGS GTNSSRESNG TMADLGNIAD NPQLSFYQLV YGLNALLLIC 840 VGVCSSGIFT KVTRKASTAL HNKLFNKVFR CPMSFFDTIP IGRILNCFAG DLEQLDQLLP 900 VGVCSGIFT KVTRKASTAL HNKLFNKVFR CPMSFFDTIP IGRILNCFAG DIEGLDQLIP 900
IFSEQFLVLS LMVIAVILIV SVLSPYILLM GAIMVICFI YYMMFKKAIG VFKRLENYSR 960
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EDRPEVI RKK PGSI FAALMA TATSSI D 75

FDRPEVLRKK PGSLFAALMA TATSSLR

_	Nucleic Acid Acc	ession #:	AA983251				
5	Coding sequence	3 :	1-1749 (un	derlined sequence	es correspond to st	art and stop codon	s)
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					GCGGCGAGTA		780
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25 .	CGGGCGGACA	AAGACGGGCC	CCGACGGCTC	GGCAGGGCTT	CATGTCTTAG	GGGTACCCAA	1020
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					TGCTGCCCCG		1140
					TGCAGCGGCC		1200
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					TCAGGCAACA		2280
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	GGGCCTATTT						2700
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					TTACTTTTAA		2940
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	GAGTGCAAGT	AATTACTATA	CTTGTAAATG	AAGATCAGTA	TTTCTGCCTA	GATCTGATAA	3060
60	AAAAATTTTC	TTGTCTTAGT	TATAAAAATT	CAAAGAAATG	TGTTACAAAG	ATACTTAGTA	3120
~	TAGCTCCTCA	GCCATAACCT	GAGACTTGGG	ATGAAATTTA	AACCAGATAC	GATTTACTTT	3180
	GCAGATCATA						3240
	GATTGTTAAG						3300
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	GTATGGTATG						3540
	TGTTTCATGA TTTTTTAATA						3600
70	TGAAAAAAA			ATTTACTTA	TTAATAAATA	TITIATGATG	3660
	* Outstand	ANNANANA.	AMMANAMA				

SEQ ID NO:194 BHB8 DNA sequence

SEQ ID NO:195 BHB8 Protein sequence

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13			APGGNRLMET NNVYVNMPTN				540	
			RQIQSPFPHT			RPFIVGITVQ	340	
	movement	TTTLUDDQTGT	MARADITIMI	HODGIGHTA				•
				. SEC	Q ID NO:196 CQA	5 DNA SEQUENCI	E	·
20	Nucleic Acid Acce	ession#:		AA088458			_	
			Co	ding sequence:		8	62-1995 (underlined sequences correspond to start and stop codons)
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45			CTGGCTGCAG					•
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UJ			TTTACAGCTT					
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			ATCCGCGAGG					
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			GCCAGGAGTT					
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SEQ ID NO:197 LBG2 DNA SEQUENCE

54-2543 (start and stop codons are underlined) Coding sequence: 5 11 21 GCGGAACACC GGCCCGCCGT CGCGGCAGCT GCTTCACCCC TCTCTCTGCA GCC<u>ATG</u>GGGC 60 GCGGAACACC GGCCCGCCTT CGCGGCAGCT GCTTCACCCC LETCTCTGCA GCCATGGGGC 00
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CCTCCGAGCC GTGCCGGCG GTCTTCAGGG AGGCTGAAGT GACCTTGGAG GCGGGAGGCG 180
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CAGCTCTGTT TAGCACTGAT AATGATGACT TCACTGTGCG GAATGGCGAG ACAGTCCAGG 300
AAAGAAGGTC ACTGAAGGAA AGGAATCCAT TGAAGATCTT CCCTGAAAAT GGCAAGGGTC 420
GAAGACACAA GAGAGTTGG GTGGTTGCTC CAATATCTGT CCCTGAAAAT GGCAAGGGTC 420 10 CCTTCCCCCA GAGACTGAAT CAGCTCAAGT CTAATAAAGA TAGAGACACC AAGATTTTCT 480 15 ACAGCATCAC GGGGCCGGGG GCAGACAGCC CCCCTGAGGG TGTCTTCGCT GTAGAGAAGG 540 AGACAGGCTG GTTGTTGTTG AATAAGCCAC TGGACCGGGA GGAGATTGCC AAGTATGAGC 600 TCTTTGGCCA CGCTGTGTCA GAGAATGGTG CCTCAGTGGA GGACCCCATG AACATCTCCA 660 TCATCGTGAC CGACCAGAAT GACCACAAGC CCAAGTTTAC CCAGGACACC TTCCGAGGGA 720 GTGTCTTAGA GGGAGTCCTA CCAGGTACTT CTGTGATGCA GGTGACAGCC ACAGATGAGG 780 20 ATGATGCCAT CTACACCTAC AATGGGGTGG TTGCTTACTC CATCCATAGC CAAGAACCAA 840 AGGACCCACA CGACCTCATG TTCACAATTC ACCGGAGCAC AGGCACCATC AGCGTCATCT 900 CCAGTGGCCT GGACCGGGAA AAAGTCCCTG AGTACACACT GACCATCCAG GCCACAGACA 960 TGGATGGGGA CGCTCCACC ACCACGGCAG TGGCAGTAGT GGAGATCCTT GATGCCAATG 1020
ACAATGCTCC CATGTTTGAC CCCCAGAAGT ACGAGGCCCA TGTGCCTGAG AATGCAGTGG 1080
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CTGAGAGCAA CCAGGGCATC CTGACAACCA GGAAGGGTTT GGATTTTGAG GCCAAAAACC 1260 25 AGCACACCCT GTACGTTGAA GTGACCAACG AGGCCCCTTT TGTGCTGAAG CTCCCAACCT 1320 CCACAGCCAC CATAGTGGTC CACGTGGAGG ATGTGAATGA GGCACCTGTG TTTGTCCCAC 1380 30 CCTCCAAAGT CGTTGAGGTC CAGGAGGGCA TCCCCACTGG GGAGCCTGTG TGTGTCTACA 1440 CTGCAGAAGA CCCTGACAAG GAGAATCAAA AGATCAGCTA CCGCATCCTG AGAGACCCAG 1500 CAGGGTGGCT AGCCATGGAC CCAGACAGTG GGCAGGTCAC AGCTGTGGGC ACCCTCGACC 1560 GTGAGGATGA GCAGTTTGTG AGGAACAACA TCTATGAAGT CATGGTCTTG GCCATGGACA 1620 ATGGAAGCCC TCCCACCACT GGCACGGGAA CCCTTCTGCT AACACTGATT GATGTCAACG 1680
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TCACAGATGA CTCAGACATC TACTGGACGG CAGAGGTCAA CGAGGAAGGT GACACAGTGG 1860 35 TCTTGTCCCT GAAGAAGTTC CTGAAGCAGG ATACATATGA CGTGCACCTT TCTCTGTCTG 1920 ACCATGGCAA CAAAGAGCAG CTGACGGTGA TCAGGGCCAC TGTGTGCGAC TGCCATGGCC 1980 40 ATGTCGAAAC CTGCCCTGGA CCCTGGAAAG GAGGTTTCAT CCTCCCTGTG CTGGGGGCTG 2040 TCCTGGCTCT GCTGTTCCTC CTGCTGGTGC TGCTTTTGTT GGTGAGAAAG AAGCGGAAGA 2100 TCAAGGAGCC CCTCCTACTC CCAGAAGATG ACACCCGTGA CAACGTCTTC TACTATGGCG 2160 AGAGGGGG TGGCGAAGAG GACCAGGACT ATGACATCAC CCAGCTCCAC CGAGGTCTGG 2220
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AGGGCAGCGG CTCCGACGCC GCGTCCCTGA GCTCCCTCAC CTCCTCCGCC TCCGACCAAG 2460 45 ACCAAGATTA CGATTATCTG AACGAGTGGG GCAGCCGCTT CAAGAAGCTG GCAGACATGT 2520 ACGGTGGCGG GGAGGACGAC TAGGCGGCCT GCCTGCAGGG CTGGGGACCA AACGTCAGGC 2580 50 CACAGAGCAT CTCCAAGGGG TCTCAGTTCC CCCTTCAGCT GAGGACTTCG GAGCTTGTCA 2640 GGAAGTGGCC GTAGCAACTT GGCGGAGACA GGCTATGAGT CTGACGTTAG AGTGGTTGCT 2700 TCCTTAGCCT TTCAGGATGG AGGAATGTGG GCAGTTTGAC TTCAGCACTG AAAACCTCTC 2760 CACCTGGGCC AGGGTTGCCT CAGAGGCCAA GTTTCCAGAA GCCTCTTACC TGCCGTAAAA 2820 55 GAAGGGTGAG GACAATCGTG TATATGTACT AGAACTTTTT TATTAAAGAA A 60 SEQ ID NO:198 LBG2 Protein sequence: CAA45177 Protein Accession #: 65 21 31 41 51 MGLPRGPLAS LLLLOVCWLO CAASEPCRAV FREAEVTLEA GGAEOEPGOA LGKVFMGCPG 60 QEPALESTION DDFTVRNGET VQERSLIKER NPLKIFPSKR ILRRIKEDWY VAPISVFENG 120 KGPFPQRLNQ LKSNKDRDTK IFYSITGPGA DSPPEGVFAV EKETGWLILN KPLDREEIAK 180 70 YELFGHAVSE NGASVEDPMN ISIIVTDOND HKPKFTQDTF RGSVLEGVLP GTSVMQVTAT 240 DEDDAIYTYN GVVAYSIHSQ EPKDPHDLMF TIHRSTGTIS VISSGLDREK VPEYTLTIQA 300 TDMDGDGSTT TAVAVVEILD ANDNAPMFDP QKYEAHVPEN AVGHEVQRLT VTDLDAPNSP 360 AWRATYLIMG GDDGDHFTTT THPESNQGIL TTRKGLDFEA KNQHTLYVEV TNEAPFVLKL 420 75 PTSTATIVVH VEDVNEAPVF VPPSKVVEVQ EGIPTGEPVC VYTAEDPDKE NQKISYRILR 480 DPAGWLAMDP DSGQVTAVGT LDREDEQFVR NNIYEVMVLA MDNGSPPTTG TGTLLLTLID 540 VNDHGPVPEP RQITICNQSP VRHVLNITDK DLSPHTSPRQ AQLTDDSDIY WTAEVNEEGD 600 TVVLSLKKFL KQDTYDVHLS LSDHGNKEQL TVIRATVCDC HGHVETCPGP WKGGFILPVL 660

GAVLALLFIL LVLLLLVRKK RKIKEPLLLP EDDTRDNVFY YGEEGGGEED QDYDITQLHR 720

X63629

Nucleic Acid Accession #:

GLEARPEVVL RNDVAPTIIP TPMYRPRPAN PDEIGNFIIE NLKAANTDPT APPYDTILLVF 780 DYEGSGSDAA SLSSLTSSAS DQDQDYDYLN EWGSRFKKLA DMYGGGEDD

5	Nucleic Acid Acce Coding sequence		NM_013	2152		DNA SEQUENCE to start and stop coo	dons)
				. (0.1000 001100	• .	,
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		TGGACTTTTT					120
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30	GTCTTAGG	AAAGCACTTC	CTAAACICIG	GATGCCTCTC	GGCCCALCCA	GGTGATGACT	1140
50	GICTIAGG						
		OBI 5 Protein seq u					
35	Protein Accession	1#:	NP_038	284			
55	1	11	21	31	41	51	
	ī	Ī .	ī	Ĩ	Ī	Ī	
		DFFYNRSNTD					60
40		NLAAADFFAG					120
40		IMRMRVHSNL WTVSNLMAFL					180 240
		CWTPGLVVLL					300
		C FSQENPERR					
45		_					
45	Shadala Salal Saa	onales de	AA569531	SE	Q ID NO:201 PAA	5 DNA SEQUENCE	
	Nucleic Acid Acco			ned sequences co	rrespond to start a	nd stop codons)	
	Com.Bood-man	•					
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50	1	GTTACTCATT	de la coccasión de la coccasió	C) COMPC MCC	THE ACCOUNT	 	60
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E E		TGGACATCCA					240
55	CAGCTATACC	TGTGGGACAT	GGGTGGTTTT	ACAATATTTA	AGAACCTGTG	GATGAGCCTC	300 360
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		GAACAATGAA					840
65		CACTCTGATA					
	SEO ID NO-202	PAA6 Protein seq	ience:				
70	Protein Accession		none found				
	1	11	21	31	41	51	
	l marchen			n policice	t. DRMCCIT.Sのな	l. Botlet ekkenn	60
75						L PCLSLSKECG K RVTETILRDF	
, 5	KOKOSSKIO	E ERRRESAGP	N LSSFWFVGN	A GRGDRPOT	WA GSKQFSG		
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SEQ ID NO: 203 PAB2 DNA SEQUENCE Nucleic Acid Accession #: XM 050197 Coding sequence: 310-1971 (underlined sequences correspond to start and stop codons) 5 31 TCACACGTGC CAAGGGGCTG GCTCAGCGGA ACCAGCCTGC ACGCGCTGGC TCCGGGTGAC 60 AGCCGCGCC CTCGGCCAGG ATCTGAGTGA TGAGACGTGT CCCCACTGAG GTGCCCCACA GCAGCAGGTG TTGAGCATGG GCTGAGAAGC TGGACCGGCA CCAAAGGGCT GGCAGAAATG 120 180 10 GGCGCCTGGC TGATTCCTAG GCAGTTGGCG GCAGCAAGGA GGAGAGGCCG CAGCTTCTGG 240 AGCAGAGCCG AGACGAAGCA GTTCTGGAGT GCCTGAACGG CCCCCTGAGC CCTACCCGCC 300 TGGCCCACTA TGGTCCAGAG GCTGTGGGTG AGCCGCCTGC TGCGGCACCG GAAAGCCCAG 360 CTCTTGCTGG TCAACCTGCT AACCTTTGGC CTGGAGGTGT GTTTGGCCGC AGGCATCACC 420 TATGTGCCGC CTCTGCTGCT GGAAGTGGGG GTAGAGGAGA AGTTCATGAC CATGGTGCTG 480 15 GGCATTGGTC CAGTGCTGGG CCTGGTCTGT GTCCCGCTCC TAGGCTCAGC CAGTGACCAC 540 TGGOGTGGAC GCTATGGCCG CCGCCGCCC TTCATCTGGG CACTGTCCTT GGGCATCCTG 600 CTGAGCCTCT TTCTCATCCC AAGGGCCGGC TGGCTAGCAG GGCTGCTGTG CCCGGATCCC 660 AGGCCCCTGG AGCTGGCACT GCTCATCCTG GGCGTGGGGC TGCTGGACTT CTGTGGCCAG 720 GTGTGCTTCA CTCCACTGGA GGCCCTGCTC TCTGACCTCT TCCGGGACCC GGACCACTGT 780 20 CGCCAGGCCT ACTCTGTCTA TGCCTTCATG ATCAGTCTTG GGGGCTGCCT GGGCTACCTC 840 CTGCCTGCCA TTGACTGGGA CACCAGTGCC CTGGCCCCCT ACCTGGGCAC CCAGGAGGAG 900 TGCCTCTTTG GCCTCCTCAC CCTCATCTTC CTCACCTGCG TAGCAGCCAC ACTGCTGGTG 960 GCTGAGGAGG CAGCGCTGGG CCCCACCGAG CCAGCAGAAG GGCTGTCGGC CCCCTCCTTG 1020 TOGOCCCACT GOTGTCCATG COGGGCCCGC TTGGCTTTCC GGAACCTGGG CGCCCTGCTT 1080 25 CONGRETOR ACCARCTOTG CTGCCGCATG CCCCGCACCC TGCGCCGGCT CTTCGTGGCT 1140 GAGCTGTGCA GCTGGATGGC ACTCATGACC TTCACGCTGT TTTACACGGA TTTCGTGGGC 1200 GAGGGGCTGT ACCAGGCGT GCCCAGAGCT GAGCCGGGCA CCGAGGCCCG GAGACACTAT 1260 GATGAAGGCG TTCGGATGGG CAGCCTGGGG CTGTTCCTGC AGTGCGCCAT CTCCCTGGTC 1320 TTCTCTCTGG TCATGGACCG GCTGGTGCAG CGATTCGGCA CTCGAGCAGT CTATTTGGCC 1380 30 AGTGTGGCAG CTTTCCCTGT GGCTGCCGGT GCCACATGCC TGTCCCACAG TGTGGCCGTG 1440 GTGACAGCTT CAGCCGCCCT CACCGGGTTC ACCTTCTCAG CCCTGCAGAT CCTGCCCTAC 1500 ACACTEGECT CCCTCTACCA CCGGGAGAAG CAGGTGTTCC TGCCCAAATA CCGAGGGGAC 1560 ACTGGAGGTG CTAGCAGTGA GGACAGCCTG ATGACCAGCT TCCTGCCAGG CCCTAAGCCT 1620 GGAGCTCCCT TCCCTAATGG ACACGTGGGT GCTGGAGGCA GTGGCCTGCT CCCACCTCCA 1680 35 CCCGCGCTCT GCGGGGCCTC TGCCTGTGAT GTCTCCGTAC GTGTGGTGGT GGGTGAGCCC 1740 ACCGAGGCCA GGGTGGTTCC GGGCCGGGGC ATCTGCCTGG ACCTCGCCAT CCTGGATAGT 1800 GCCTTCCTGC TGTCCCAGGT GGCCCCATCC CTGTTTATGG GCTCCATTGT CCAGCTCAGC 1860 CAGTCTGTCA CTGCCTATAT GGTGTCTGCC GCAGGCCTGG GTCTGGTCGC CATTTACTTT 1920 GCTACACAGG TAGTATTTGA CAAGAGCGAC TTGGCCAAAT ACTCAGCG<u>TA G</u>AAAACTTCC 1980 40 AGCACATTGG GGTGGAGGGC CTGCCTCACT GGGTCCCAGC TCCCCGCTCC TGTTAGCCCC 2040 ATGGGGCTGC CGGGCTGGCC GCCAGTTTCT GTTGCTGCCA AAGTAATGTG GCTCTCTGCT 2100 GCCACCCTGT GCTGCTGAGG TGCGTAGCTG CACAGCTGGG GGCTGGGGCG TCCCTCTCCT 2160 CTCTCCCCAG TCTCTAGGGC TGCCTGACTG GAGGCCTTCC AAGGGGGTTT CAGTCTGGAC 2220 TTATACAGGG AGGCCAGAAG GGCTCCATGC ACTGGAATGC GGGGACTCTG CAGGTGGATT 2280 45 ACCCAGGCTC AGGGTTAACA GCTAGCCTCC TAGTTGAGAC ACACCTAGAG AAGGGTTTTT 2340 GGGAGCTGAA TAAACTCAGT CACCTGGTTT CCCATCTCTA AGCCCCTTAA CCTGCAGCTT 2400 CGTTTAATGT AGCTCTTGCA TGGGAGTTTC TAGGATGAAA CACTCCTCCA TGGGATTTGA ACATATGAAA GTTATTTGTA GGGGAAGAGT CCTGAGGGGC AACACAAAG AACCAGGTCC 2460 2520 CCTCAGCCCC ACAGGCACTG GTCTTTTTTG CTNGANTCCA CCCCCCCCT CTTTACCCTT 50 SEQ ID NO:204 PAB2 Protein sequence: Protein Accession #: XP 050197 55 11 21 31 51 MVORLWYSKL LRHRKAQLLL VNLLTFGLEV CLAAGITYVP PLLLEVGVEE KFMTMVLGIG 60 PVLGLVCVPL LGSASDHWRG RYGRRRPFIW ALSLGILLSL FLIPRAGWLA GLLCPDPRPL 120 ELALLILGVG LLDFCGQVCF TPLEALLSDL FRDPDHCRQA YSVYAFMISL GGCLGYLLPA 180 IDWDTSALAP YLGTQEECLF GLLTLIFLTC VAATLLVAEE AALGPTEPAE GLSAPSLSPH 240 CCPCRARLAF RNLGALLPRL HQLCCRMPRT LRRLFVAELC SWMALMTFTL FYTDFVGEGL 300 YQGVPRAEPG TEARRHYDEG VRMGSLGLFL QCAISLVFSL VMDRLVQRFG TRAVYLASVA 360 65 AFPVAAGATC LSHSVAVVTA SAALTGFTFS ALQILPYTLA SLYHREKQVF LPKYRGDTGG 420 ASSEDSLMTS FLPGPKPGAP FPNGHVGAGG SGLLPPPPPAL CGASACDVSV RVVVGEPTEA 480 RVVPGRGICL DLAILDSAFL LSQVAPSLFM GSIVQLSQSV TAYMVSAAGL GLVATYPATQ 540 VVFDKSDLAK YSA 70 SEQ ID NO:205 PAJ3 DNA SEQUENCE Nucleic Acid Accession #: AK002126 1-1593 (underlined sequences correspond to start and stop codons) Coding sequence: 75 11 21 31 51 . ATGGTTCGCC GGGGCTGCT TGCGTGGATT TCCCGGGTGG TGGTTTTGCT GGTGCTCCTC

TGCTGTGCTA TCTCTGTCCT GTACATGTTG GCCTGCACCC CAAAAGGTGA CGAGGAGCAG

CTGGCACTGC CCAGGGCCAA CAGCCCCACG GGGAAGGAGG GGTACCAGGC CGTCCTTCAG

GAGTGGGAGG AGCAGCACCG CAACTACGTG AGCAGCCTGA AGCGGCAGAT CGCACAGCTC

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		TGCAGGAGAG TGGGTCTGGA					300 360
5	TATGCAGCAG CTTACCCGCC	TGCCTTTCGA ACCCCGAGGA	TAGCTTTACT GAAGCCTGTG	CTACAGAAGG AGGAAGGACA	TGTACCAGCT AGCGGGATGA	GGCCACAGAG GGAGACTGGC GTTGGTGGAA CAATCACCGT	420 480 540 600
	CCTTACACGG TTGTATGAGC	CCTCTGATTT	CATAGAAGGG AGGGGACCAC	ATCTACCGAA AAACACGAAT	CAGAAAGGGA TCAAACGGCT	CAAAGGGACA CATCTTATTT	660 720 780
10	TTCAGGGAGA AAAGAAGAAA	TGTGCATTGA TAAATGAAGT	GCAGGATGGG CAAAGGAATA	AGAGTCCATC CTTGAAAACA	TCACTGTTGT CTTCCAAAGC	CATGCAGAAT TTACTTTGGG TGCCAACTTC ACTTGATGTT	840 900 960 1020
15	GGAGCCCGCT TTCACATCTG TATCCAGTTC CCTCCCTTGG	TCTGGAAGGG AATTCCTCAA TTTTCAGTCA AACAGCAGCT	AAGCAACGTC TACGTGTAGG GTACAATCCT GGTCATAAAG	CTTCTCTTTT CTGAATACAC GGCATAATAT AAGGAAACTG	TCTGTGATGT AGCCAGGGAA ACGGCCACCA GATTTTGGAG	GGACATCTAC GAAGGTATTT TGATGCAGTC AGACTTTGGA	1080 1140 1200 1260
20	ATCAAAGGCT ATAGTGGTAC GACGAGCTGA TCCCACGGCC	- GGGGCGGAGA GGACGCCTGT CCCCCGAGCA AGCTGGGCAT	GGATGTGCAC GCGAGGACTC GTACAAGATG GCTGGTGTTC	CTTTATCGCA TTCCACCTCT TGCATGCAGT AGGCACGAGA	AGTATCTCCA GGCATGAGAA CCAAGGCCAT	TGATCTGGAC CAGCAACCTC GCGCTGCATG GAACGAGGCA CCTTCGCAAA	1320 1380 1440 1500 1560
25	CAGAAACAGA	AGACAAGTAG	CAAAAAAACA	TGA			
25	SEQ ID NO:206 P Protein Accession		nce: NP_060841				
30	1	11 :	21 :	31 I	41 i	51 I	
50	mvrrgllawi : Eweeqhrnyv : Flhsqvdkae : Aiesaletin :	SSLKRQIAQL I VNAGVKLATE I	KEELQERSEQ 1 YAAVPFDSFT 1	LRNGQYQASD : LQKVYQLETG :	AAGLGLDRSP LTRHPEEKPV	PEKTQADLLA RKDKRDELVE	60 120 180 240
35	RPFGPIMKVK I KEEINEVKGI I FTSEFLNTCR I FGMTCQYRSD I	neklnmantl Lentskaanf Lntqpgkkvf	INVIVPLAKR ' RNFTFIQLNG ' YPVLFSQYNP (VDKFRQFMQN EFSRGKGLDV GIIYGHHDAV	FREMCIEQDG GARFWKGSNV PPLEQQLVIK	RVHLTVVYFG LLFFCDVDIY KETGFWRDFG	300 360 420 480
40	DELTPEQYKM						100
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	Nucleic Acid Acces Coding sequence:	ssion #:	AF189723 1-2712 (underlin		1D NO:207 PAJ5 respond to start a		
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45 50	Coding sequence: 1 ATGATTCCTG	11 TATTGACATC CTGATCTTCA	1-2712 (underlin 21 AAAAAAAGCA GAATGGTCTA	31 AGTGAATTAC	TESPOND TO START AT THE START AT THE START AS A SAGTTAGTCA A A SAGTTAGTCA A SAGTTAGTAGTCA A SAGTTAGTCA A SAGTTAGTAGTAGTAGTAGTAGTAGTAGTAGTAGTAGTAGT	51 AGTTGCAAGC TAGGCGAGCC	60 120
	1 ATCATTCCTG ATTCATGGCT TCTCAGGTTTA	11 TATTGACATC CTGATCTTCA GGAATGAGTT AAAATCCCCT	1-2712 (underling 21 AAAAAAAGCA GAATGGTCTA TGATATTAGT TATTATGCTG	31 AGTGAATTAC AACAAATGTG GAAGATGAGC CTTCTGGCTT	41 CAGTCAGTGA AAGTTAGTCA CACTGTGGAA CTGCAGTCAT	51 AGTTGCAAGC TAGGCGAGCC GAAGTATATT CAGTGTTTA	120 180 240
50	1 ATGATTCCTG ATTCTCAAG TTTCATGGCT TCTCAGTTTA ATGCATCAGT	11 TATTGACATC CTGATCTTCA GGAATGAGTT	1-2712 (underling the control of the	31 	41 CAGTCAGTGA AAGTTAGTCA CACTGTGGAA CTGCAGTCAT TACTTATCGT	51 AGTTGCAAGC TAGGCGAGCC GAAGTATATT CAGTGTTTTA	120 180
	1 ATGATTCCTG ATTICATGGCT TCTCAGTTTA ATGCATCAGT GCCTTTGTTC CCAGAATGCC	11 TATTGACATC CTGATCTTCA GGAATGAGTT AAAATCCCCT TTGATGATCG AGGAATATCG ATTGTGTGCG	1-2712 (underling) 21 AAAAAAAGCA GAATGGTCTA TGATATTAGT TATTATGCTG CGTCAGTATC TTCAGAAAAA TGAAGGAAAA	31 AGTGAATTAC AACAAATGTG GAAGATGAGC CTTCTGGCTT ACTGTGGCAT TCTCTTGAAG	41 CAGTCAGTGA AAGTTAGTCA CACTGTGGAA CTGCAGTCAT TACTTATCAT AATTGAGTAA CACTTGCCCG	51 AGTTGCAAGC TAGGCGAGCC GAAGTATATT CAGTGTTTTA TGTTACAGTT ACTTGTGCCA AGACTTGGTT	120 180 240 300 360 420
50	1	11 TATTGACATC CTGATCTTCA GGAATGAGTT TANAATCCCCT TTGATGATGC AGGAATATCG ATTGTGTGCCC ATCTTTCCAT	1-2712 (underling) 21	31 AGTGAATTAC AACAAATGTG GAAGATGAGC CTTCTGGCTT ACTGTGGCAA TCTCTGAAG TCTGTGAGCATA AGATAGAGTTC AGCTTGACAG	41	51 AGTTGCAAGC TAGGCGAGCC GAAGTATATT CAGTGTTTTA TGTTACAGTT ACTTGTGCCA AGACTTGGTT ACCTTGTTTT GCCTTGTTTT	120 180 240 300 360 420 480 540
50 55	1	11 TATTGACATC CTGATCTTCA GGAATGAGTT AAAATCCCCT AGGAATATCG ATTGTGTGCC ATTGTGTGCCT ATCTTTCCAT CTCCTCAGCC	1-2712 (underling the control of the	31 AGTGAATTAC AACAAATGTG GAAGATGAGC CTTCTGGCTT ACTGTGGCAA TCTCTTGAAG TCTCTTGAAG TTGGAGCATA GATAGAGTTC AGCTTGACAG AATGGAGATC	41 CAGTCAGTGA AAGTTAGTCA CACTGTGGAA CTGCAGTCAT TACTTATCGT AATTGAGTAA CACTTGCCCG CTGCTGACTT GTGAGACAAC TTGCATCGAG	51 AGTTGCAAGC TAGGCGAGCC GAAGTATATT CAGTGTTTTA TGTTACAGTT ACTTGTGCA AGACTTGTTT ACGCTTGTTT ACGCTTGTTT ACGCTTGTTT ACGCTTGTTT AAGTAACATT	120 180 240 300 360 420 480 540 600
50	1	11 TATTGACATC CTGATCTTCA GGAATGAGTT AAAATCCCT TTGATGATGC AGGAATATCG ATTGTGTGCG CAGTTTCCAT CTCCTCAGCC GAACACTGGT AATTTGGGGA	1-2712 (underling the control of the	31 AGTGAATTAC AACAAATGTG GAAGATGAGC CTTCTGGCTT ACTGTGGCAA TCTCTTGAAG TTGGAGCATT AGATGAGGTTC AGCTTGACAG AATGAGGTTC AAGCAAAGG ATGATGCAAAG	41	51 AGTTGCAAGC TAGGCGAGCC GARGTATATT CAGTGTTTTA TGTTACAGTT ACTTGTGCTA AGACTTGTTT ACGCTTGTTT ACGCTTGTTT TGCAACAGGA ACCAACAGGA	120 180 240 300 360 420 480 540 600 660 720
50 55	1	11 TATTGACATC CTGATCTTCA GGAATGAGTT AAAATCCCCT TTGATGATGC AGGAATATCG ATTGTGTGCG CACTTTGCCT ATCTTTTCCAT CTCCTCAGCC GAACACTGGT	1-2712 (underling the control of the	31 AGTGAATTAC AACAAATGTG GAAGATGAGC CTTCTGGCTA ACTGTGGCAA TCTCTGAGC TTGGAGCATA GATAGAGTTC AGCTTGACAG AATGGAGATC AAACCAAAG	41 CAGTCAGTGA AAGTTAGTCA CACTGTGGAA CTGCAGTCAT TACTTATCGT AATTGAGTAA CACTTGCCG GTGCAGTTGCTG GTGAGACAAC GTGTTGTCAG GTGTTGTCAG GTGTTGTCAG CAGAAGAGGC CCTTTTACTC	51 AGTTGCAAGC TAGGCGAGCC GAAGTATATT CAGTGTTTA ACTTGCCA AGACTTGGTT ACCCTTGTTCT AAGTAACATT TGGAACAGGA ACCAAAAACC CTTTGGTATA	120 180 240 300 360 420 480 540 600 660
50 55	1	11 TATTGACATC CTGATCTTCA GGAATGAGTT AAAATCCCT TTGATGATGG ATTGTGTGCG CAGTTTGCCT ATCTTTCCAT CTCCTCAGCC GAACACTGGT AATTTGGGGA AGAGCATGGG TCATGTTGGGT GTTTGGGT GTTTGGCTGTTTTGGTGGT	1-2712 (underling the control of the	31 AGTGAATTAC AACAAATGTG GAAGATGAGC TTCTGGCTT ACTGTGGCAT ACTATGAGGTTC AGTTGACAG AATGAGGTC AATGAGATC AATGAGATC AAACAACTT AACAACAACTTT AACGAAAAG CCTGAAGGTC	41 CAGTCAGTGA AAGTTAGTCA CACTGTGGAA CTCAGTCAT TACTTATCGT TACTTGAGTAA CACTTGCCCG CTGCTGACTT GTGAGACAAC TTGCATCAGT GTGAGACAAC GTGTTGTCAT CAGAAGAGGC CCTTTTTACT ATATCCTGGA TCCCATTGT	51 AGTTGCAAGC TAGGCGAGCC GAAGTATATT CAGTGTTTTA ACTTGTCCA AGACTTGGTT ACCCTTGTTC AAGTAACAT TCGAACAGGA ACCAAAAACC CTTTGGTATA AATGTTACA GGTCACAGTA GGTCACAGTA	120 180 240 300 360 420 480 540 660 720 780 840 900
50 55	1	11 TATTGACATC CTGATCATCA GGAATGAGTT AAAATCCCT TTGATGATGGG ATTGTGTGCG ATTGTTTCCAT CTCCTCAGCC GAACACTGGT AATTTGGGA ACACTGGT ATTTTGGGA ACACTGGT TTGGTGTTGTTTTGTT	1-2712 (underling the control of the	31 AGTGAATTAC AACAAATGTG GAAGATGAGC CTTCTGGCTT ACTGTGGCAA TCTCTTGAAG TTGGAGCATT AGCTTGACAG AATGAGATTC AGCTTGACAG AATGAGATTC AAAGCAAAGG ATGATGCAAG AAACAACTTT CTGGGAAAAG CCTGAAAGG AACAAACGTAAAGGTAAAAGGGTC AAGAAAAGGGTC AAGAAAAGGGTC AAGAAAAGGGTC AAGAAAAGGGTC	41 CAGTCAGTGA AAGTTAGTCA CACTGAGTCAT TACTTATCGT AATTGAGTAA CACTTGCCG CTGCTGACTT GTGAGACAAC TTGCATCGAG TTGCATCGAG CAGTAGTTGTCAT CAGAAGAGGC CCTTTTACTC ATATCCTGA TCCCATTGT CCACTTGTA	51 AGTTGCAAGC TAGGCGAGCC GAAGTATATT CAGTGTTTTA ACTTGTCCA AGACTTGGTT ACCCTTGTTC AAGTAACAT TCGAACAGGA ACCAAAAACC CTTTGGTATA AATGTTACA GGTCACAGTA GGTCACAGTA	120 180 240 300 360 420 480 540 660 720 780 840 900 960
50 55 60	1	11 TATTGACATC CTGATCTTCA GGAATGAGTT TAGATGATCC TTGATGATCC ATTGTGTGCG ATTGTCAT CTCCTCAGCC GAACACTGGT AACATTTGGGGA ACACTGGT TCATGTGTGTT TTGGTGTATC TTCGTGTTAC TCTCTGGCTT TCGTGTTAC TCTCTGCCTTAC TCATGTGTTAC	1-2712 (underling the control of the	31 AGTGAATTAC AACAAATGTG GAAGATGAGC CTTCTTGGCTA TCTCTTGAAG TTGGAGCATA ACTGTGACATA AACAAATGT AATGAAGATC AAACAAATGT CTGGAGAAAAG ATGAAGAAAGC AAGAAAAGG AAGAAAAGC AAGAAAAGC AAGAAAAGC AAGAAAAGC AAGAAAAGC ATTGAAAAAGC ACTGAAAAGG	41 CAGTCAGTGA AAGTTAGTCA CAGTGAGTCAT TACTTATCGT AATTGAGTA CACTTGCCG CTGCTGACTT GTGAGACAAC CTGCAGTCAT CAGTGACAT CAGTGACAT CAGAGAGAG CCTTTTTACTC ATATCCTGGA TCCCCATTGT CCATTGTGA ATAAACTGG GTCTGCATG	51 51 AGTTGCAAGC TAGGCGAGCC GAAGTATATT ACTTGTCCA AGACTGGTT ACGCTTGTTC AAGACACAGA ACCAAAAACC CTTTGGTATA AATGTTTACT GGACAGGGA ACCAAAAAC ATGGTATAC AAAGCTGCT AAAGCTGCT AAAGCTGCT TAGGACAGGC TAGACAGCC TGAGGTTACC TGAGGTTACC	120 180 240 300 360 420 480 540 600 720 780 840 900 960 1020
50 55 60	1	11 TATTGACATC CTGATCTTCA GGAATGAGTT TAAAATCCCCT TTGATGATGC AGGAATATCG ATCTTTGCAT CTCCTCAGCC GAACACTGGT AATTTGGGA AGACATGGT TTGGTTTGGT	1-2712 (underling the control of the	31 AGTGAATTAC AACAAATGTG GAAGATGAGC CTTCTGGCTA ACTGTGGCAA TCTCTTGAAG TTGAGCATTA AGCTTGACAG AATGGAGATTC AGCTTGACAG AATGAGATTC TGGGAAAAG AACAACTTT CTGGGAAAAG CATGAAGGTC AAGAAAAGGGAAAAAGGGAAAAAAGGGAAAAGGGTC ATGTTCAGAAGGT ATTGTTCAG ACTTCAGATG ATTGTTCATG CCGGCCTGTG	41 CAGTCAGTGA AAGTTAGTCA CACTGAGGAT CACTGAGGAT AATTGAGTAA CACTTGCCG CTGCTGACTT GTGAGACAAC TTGCATCAGAG GTGTTGTCAT CAGAAGAGGC CCTTTTACTC ATATCCTGA TCCCATTGT CCATTGTGAA ATAAAACTGG GTCTTGCATGAG GTCTTGCATGAG ATAAAACTGG GTCTTGCATGT TGTGCAATGA	51 AGTTGCAAGC TAGGCGAGCC GAAGTATATT CAGTGTTTA ACTTGTCA AGACTTGTT ACGCTTGTTT ACGCTTGTTT ACGCTTGTTT ACGCAACAGGA ACCAAAAACC CTTTGGTATA AATGTTTACT GGTCACAGTG AAAGCTGCCT AACACTGACT TCACGGATTC TCATGGATTA TCATGGATTC	120 180 240 300 360 420 480 540 660 720 780 960 1020 1020 1140 1200
50 55 60	1	11 TATTGACATC CTGATCTTCA GGAATGAGTT AAAATCCCCT AGGAATATCG ATGTGTGCC ATGTTTCCAT CTCCTCAGCC GAACACTGGT CTCTCAGCG AACACTGGT TCATTGGGGA ACACTGGT TCATTGGCTGT TCGTGTGTTT CTCTCAGCCTGT TCGTGTGTTT CTCTCAGCCTG TCACTGTTACCAG ATAATCAATT CTCTTAGCAG CTCTAATGGG CTCTAATGGA CTCTAATGA CTCTAATGAATGA CTCTAATGAATGAATGAATAATAATAATAATAATAATAATAAT	1-2712 (underling the control of the	31 AGTGAATTAC AACAAATGTG GAAGATGAGC CTTCTGGCTA TCTCTGGCAA TCTCTGAGG TTGGAGCATA GATAGAGTTC AAGCAAAGG AATGGAGATC CTGGGAAAAG CATAGAGTTC CTGGGAAAAG ATGCAAGG ATGTTCTGAGGTTC AAGAAAGGG ATTTCTTGATG ACTTCAGATG ATTCTTGATG ACTTCAGATG CCGGGCTGTG GCAGGGCCT	41 CAGTCAGTGA AAGTTAGTCA CAGTGAGTCAT TAGTTATCGT AATTGAGTAA CACTTGCCGG CTGCTGACTT GTGAGACAAC CTGCAGTCAT TTGCATCGAG CTGTTGTCAT CAGAAGAGGC CCTTTTACTC ATATCCTGAA ATAAACTGG GTCTGCATGT CCATTGT CCATTGTC ATATCCTGAA ATAAACTGG GTCTGCATGT TTGTCCATGC TTGTGCATGC TTGTGCAATGA TAATTGCTCT	51 AGTTGCAAGC TAGGCGAGCC GAAGTATATT CAGTGTTTTA TGTTACAGTT ACGTTGTTCT ACGTTGTTCT ACGTAGACACGA ACCAAAAACC CTTTGGTATA AATGTTTACT GGTCACAGTG AAAGCTGCTT AAAGCTGCTT AAAGCTGCTT AAAGCTGCCT AAAACC TTAGTACA TGTACAGTG AAAACC TTAGTACA TGTACAGTTC TCATGACTTC TCATGGATTC TCATGGATTC	120 180 240 360 420 480 540 660 720 780 840 900 900 1020 1080
50 55 60 65	1	11 TATTGACATC CTGATCTTCA GGAATGAGTT AAAATCCCT TTGATGATGC ACTTTGCCT ATCTTTCCAT ATCTTTCCAT ATCTTTCCAT ATCTTTCCAT ATCTTTCCAT ATCTTTCCAT ATCTTTGGGGA AGAGCATGGT TCGTGTTAT CTCTTGGCTG TTGGTGTTAT CTCTTAGCAG ATAATCAATT CTCTTAGCAG CTCTAATGG ATGGACTTCA GGATGGCTGT GGATGGCTGT GGATGGCTGT GGATGGCTGT ATGGACTTCA GGATGGCTGT GGATGGCTGT	1-2712 (underling the control of the	31 AGTGAATTAC AACAAATGTG GAAGATGAG CTTCTTGGCTT ACTGTGGCAA TCTCTTGACA TGAGAGATTAC AAGAAGAGATTC AGCTTGACAG AATGAGATTC AGCTTGACAG AATGAGATTC AAGCAAAGG ATGATGCAAG AAACAACTTT CTGGGAAAAG AAACAACTTTCATCAGATC AAGAAAAGGGCATTCAGATC ATTGTTGATG ACTGCAACAC ACTGCAACAC CACCGAACAC CACCGAACAC	41 CAGTCAGTGA AAGTTAGTCA CACTGTGGAA CACTGTGGAA CACTTGCCG CTGCTGACTT GTGAGACAAC TTGCATCGAG GTGTTGCATCGAG GTGTTGCATCGAG ATATCCTGGA TCCCATTGTCAT ATATCCTGGA TCCCATTGTGAA ATAAAACTGG GTCTGCATGT GTGCATGC GTGATGTTGCATGAATAACTCG AGAAGAGACAC AGAAGACAC AGCAGGACAG	51 AGTTGCAAGC TAGGCGAGCC GARGTATATT CAGTGTTTTA TGTTACAGTT ACTTGTGCTA AGACTTGTTT ACGTTGTTCT AAGTACAGT ACACTAGATA ACACTTGTTC AAGTACAGT ACACAGA ACCAAAAACC CTTGGTATA AATGTTACT AAGCTGCCT AAAGCTGCCT AAAGCTGCCT TAGGATTAC TCATGGATTC TCATGGATTC TCCATGAATT TCCAATGAAT TTTAGCTTTA	120 180 240 360 420 480 660 720 780 840 900 960 1020 1140 1260 1320 1380
50 55 60 65	1 ATGATTCCTG ATTCTCCAAG TTTCATGGCT TCTCAGTTTA ATGCATTGTC CCAGATGCC CCAGGTGATA GAGGCTGTGG AAGGTGACAG GCCTTTATG GAAATCCTG AAAGTGACAG ATTAGTGTAA ATTAGTGTAA AAGGATGACAC ATTAGTGTAA AAGGATGACAT ATTAGTGTAA AAGGATGACAT ATTGTTGAAA AAGATGAAA AAGAATGAAA AAGAACATTA ATGGGTCTTG GAGCAAAAGT TGTTTTATGG GGGCAGACCT	11 TATTGACATC CTGATCTTCA GGAATGAGTT AAAATCCCCT AGGAATATCG ATTGTGTGCC ATTTTCCAT CTCCTCAGCC GAACACTGGT CTCTCAGCG AACACTGGT TTGGTGTGTT TTGGTGTTTTCCTT TTGGTGTTTTCCTT TTGGTGTTTAC ATTATCAGC CTCTAATCAGC CTCTAATCAG ATGACTTTA CAGTGTTTT CTCTTAGCAG ATGACTTTA TGACACTTAT TGACACTTTA TGACACTTTA TGACACTTTA TGACACTTTA TGACACTTTA TGACACTTTA	1-2712 (underling the control of the	31 ACTGAATTAC AACAAATGTG GAAGATGAGC CTTCTGGCTA TCTCTGGCAA TCTCTGGCAA TCTCTTGAGG TTGGAGCATA AATGAAGTTC AAGAAAAGG AATGAAGAC AAACAAACTTT CTGGGAAAAG AAACAAACTTT CTGGGAAAAG AATTCAGAAAAGG ATTTCAGATG ACTTCAGATG ACTTCAGATG ACTTCAGATG ATTTCTTGATG GCGGCCTTG GCAGCGCCT ATCAGAAAG CACCGAACAC ATTAAGTACT AGAGATGTGT	41 CAGTCAGTGA AAGTTAGTCA CAGTGAGTCAT TAGTTATCGT AATTGAGTAA CACTTGCCGG CTGCTGACTT GTGAGACAAC CTGCAGTCAT TTGCATCGAG CTGCTTGTCAT CAGAAGAGGC CCTTTTACTC ATATCCTGAA ATAAACTGG GTCTGCATGT TCTGCATGT TCTGCATGT TTGTGCATGT TAGTGATACAC TAATTGCTCT TGTGCAAGACAGC TGAATACCC GTGATACCC GTGATACCC GTGATACCC GTGATACCAC AGCACGACAC GTACTACATA ACCAACAAGA	51 AGTTGCAAGC TAGGCGAGCC GAAGTATATT CAGTGTTTTA TGTTACAGTT ACTTGTGCCA AGACTTGTTT ACGTTGTTT ACGTTGTTT AGGACAGGA ACACAGAAAAC CTTTGGTATA AATGTTTACT GGTACAGTA AAAGCTGCCT AACACTGACG TCAGGGTTACT TCATGGATAT TCAATGATT TCCAATGATT TCCAATGATT TCCAATGAGT TCATGGATT TCCAATGAGT TCATGGATT TCCAATGAGT TCATGATT TCCAAGAGAAA CAGAGAAA GAAGGCACCC	120 180 240 300 360 420 480 540 660 720 900 900 900 1020 1140 1200 1260 1320
50 55 60 65 70	1	11 TATTGACATC CTGATCTTCA GGAATGAGTT TAGATGATCC TTGATGATCC TTGATGATCC ATTGTGCC ATTGTCCAT CTCCTCAGCC GAACACTGGT AACATTGGGA ACACCATGGA TCATGTGGT TTGGTGTTA CTCTTAGCAT CTCTTAGCAT CTCTTAGCAT ATAATCAATT CTGTTAGTAT CTGTTAGCAT ATAATCAATT CTGTTAGCAT ATAATCAATT CTGTTAGCAT ATGACTGTTA GAAGGCCTTCA GGATGGCTGT AAGGTGCTTTA AAGGTGCTTA TGACACTTAC CGGGACTCAG	1-2712 (underling the control of the	31 AGTGAATTAC AACAAATGTG GAAGATGAGC CTTCTGGCTA TCTCTTGAGG TTGGCAAATGAGATTGACAAAAGG AATGGAGATC AAACAAAAGG AATGAAAAGG AATGAAAAGG AATGAAAAGG AATGAAAAGG AATGAAAAGG ACTTCAGATG ACTTCAGATAGTAC ACTTCAGATAGTAC ACTTCAGAAAAG CACCGAACAC ATTAAGTACT TTGGCTTCTG	41 CAGTCAGTGA AAGTTAGTCA CAGTGAGTCA CACTGTGGAA CTGCAGTCAT TACTTATCGT AATTGAGTAA CACTTGCCG CTGCTGACTT GTGAGACAAC CAGTGAGCAT CAGTTGTCAT CAGAAGAGG CCTTTTACTC ATATCCTGA TCCCATTGT CCATTGTGA TCCATTGTGATCATCGT TGTGATGTGT TGTGCATTGT TGTGCATTGT TGTGCATTGT TGTGCATGC CTGATGTTGT CTGAATACCC AGCAGGACAG GTACTACATA GTCCTGAACT GTACTACATA GTCCTGAACT	51 AGTTGCAAGC TAGGCGAGCC GAAGTATATT ACTTGTACAGT ACTTGTGCA AGACTTGTTT ACGTTGTTT ACGTTGTTT ACGTTGTTT ACGTACAGT ACACTGGTT TGGAACAGGA ACCAAAAAC CTTTGGTATA AATGTTTACT GGTCACAGTG TAAGGTTCCT TCATGGATTC TCATGGATTC TCATGGATTC TCATGGATTC TCATGGATTC TCATGGATTC TCATGGATTC TCATGAGTTC TCATGAGTTC TCATGAGTT CCAGAGCAAA ACCTGACC GGAGCACGC GGGACAGCTG GAGGCACCC GGGACAGCTG	120 180 240 300 360 420 480 660 720 840 900 900 1020 1080 1140 1260 1320 1380 1440 1560
50 55 60 65	1	11 TATTGACATC CTGATCTTCA GGAATGAGTT AAAATCCCCT TTGATGATGCC AGGAATATCG ATTGTGTGCC CAGTTTGCCT ATCTTCCAT CTCCTCAGCC GAACACTGGT GTTTGGCTG TTGGTTGGTTAT CTCTGGGCTG TTGGTGTTAT CTCTGGCTG ATAATCAATT CTCTTAGCAG ATAATCAATT CTGTTAGCAG ATAATCAATT CTGTTAGCAG CTCTAATGG ATGACTTTA GGATGGCTTT TGACACTTTA CGGGACTCAA GGATGGCTTT TGACACTTAC CGGGACTCAG GCTTGGTGGG TTGCCTCAGG	1-2712 (underling land) 21 AARARAAGCA GRATGGTCTA TGATATTAGGT TTATATGGT TTCAGARAAA TCAGAGAAAA TCAGAGACAC AGCTGCAACT CAGATGTGG GGTTTTTAGGA TGCTGCAGTTT TGGGAAGCTAC TGAGAAGCTAC CAGATGTGG CTGTAATGT TGAGAATGTG CAGATGTTG CAGATGTTG CAGATGTTG CAGATGTTG CAGATGTT TGAGAATGTT TGAGAATGTT TGAGAATGTT TGAGAATGTT TGAGAATGTT TGAGAATGTT TGAGAAGCTAC ACAAGACTAC TAAGTGTTA TCAGAGACTAC TAAGTGTTA TCAGAGAGTA AGTATCTTTGT AGTATCATTA AGTATCAATA	31 AGTGAATTAC AACAAATGTG GAAGATGAGC CTTCTGGCTA ACTGGCTA ACTGGCTA ACTGGGCTA ACTGGGCAA TTGGGGAGTTC AGCTTGACAG AATGGAGTTC AGCTTGACAG AATGGAGATC CTGGGAAAAG AACAAACG ATTCAGAAAG ATTTCATTGATG GCGGGCCTTG GAAGGGGCCT ATCAGAAAG CAACAGAACA ATTAAGTACT AGAAAGC ATTAGGTTCT CAGCAACAC ATTAAGTACT AGAGATGTT TTGGCTTCTG CCACCTAGAA AAAATGATTA AAAATGATTA	41 CAGTCAGTGA AAGTTAGTCA CACTGTGGAA CACTGTGGAA CACTTGCCGG CTGCTGCCGG CTGCTGCAGT GTGAGACAAC CTGCAGTCAT CAGAAGAGGC CCTTTTACTC ATATCCTGAA ATAAAACTGG GTCTGCATGC CTGCATGT CTGCATGAATACC CTGAATACCT CTGAATACCT CTGAATACCT CTGCAGCACG GTCCTGACT CTGCACACACACAC CTCCTGACT CTGGTGTGAA CCTGGAGATTC CTGGTGTGAA CTGCAGGACTC CTGGTGTGAA CTGCAGGACTC	51 AGTTGCAAGC TAGGCGAGCC GAAGTATATT CAGTGTTTTA ACTTGTCCA AGACTTGTTT ACCTTGTCT AAGTAACATT TGGAACAGG ACAAAAACC CTTTGGTATA AATGTTTACT GATCACAGG AAGACTAGTT TCAGACAGGT TAACATTACT TCATGACAGG TAACATT TCAGATACAT TCATGATTAC TCATGATTAC TCATGATTAC TCATGATTAC TCATGAATT TCCAATGAAT TTTTAGCTCT ACCAGAGATT CCAGAGCATA GAAGGCACGC GGGACAGCTG AGAAGCTGTT ACAGGAGCACT ACAGGAGACTT ACAGGAGACTT ACAGGAGACACT ACAGGAGACT ACAGGAGACTT ACAGGAGACTT ACAGGAGACTT ACAGGAGACTT ACAGGAGACTT ACAGGAGACTT ACAGGAGACTT ACAGGAGACT	120 180 240 300 480 540 600 660 780 840 900 1020 1180 1260 1380 1440 1500 1560 1680
50 55 60 65 70	1	11 TATTGACATC CTGATCTTCA GGAATGAGTT ARAATCCCT AGGAATATCG ATTGTGCC AGGAATATCG ATTTTCCAT CTCCTCAGCC GAACACTGGT CTCTCAGCG AACACTGGT TTGGTGTGT TTGGTGTGTT TTGGTGTTTC CTCTAGCGC GAACACTGTT CTCTTGGCTGT TTGGTGTTAC ATTATCAGT CTCTTAGCAG ATGATCTCA GGATGGCTTA TGACACTTAC GGATGGCTTA TGACACTTAC CGGGACTCAG CTTGGTGGG TTGCCTCAGG TTCCCTCAGG TTCCCTCAGG TCCCCAGTCC	1-2712 (underling the control of the	31 AGTGAATTAC AACAAATGTG GAAGATGAGC CTTCTGGCTA TCTCTGGCAA TCTCTGGCAA TCTCTTGAGG TTGGAGCATA GATAGAGTTC AAGAAAAGG AATGGAGATC CTGAAGGTC CTGAAGGTC AAGAAAAGG ATTCTTGTCAG ACTTCAGATG ATTCTTGATG ACTTCAGATG ACTTCAGATG ACTTCAGATG ACTTCAGATG ACTTCAGATG ACTTCAGATG ACTTCAGATG CCCGACACA CACCGAACAC TTGAGATGTT TTGGCTTCTG CCACCTAGAA AAAATGATTA TATTCCAAAA	41 CAGTCAGTGA AAGTTAGTCA CAGTCAGTGAA CTGCAGTCAT TACTTATCGT AATTGAGTAA CACTTGCCGG CTGCTGACTT GTGAGACAAC CTGCAGTCT CTGCAGTCAT TTGCATCGGA ATACTTGCAT ATACTTGAT ATACTTGAT ATACTTGAT ATACTTGAA ATAAAACTGG GTCTGCATGT TGTGCATGA TAGTGTATA TAGTGTATA CCGAGGACAG GTACTACATA ACCAACAAGA GTCCTGAACT CTGGTGTGAACT CTGGTGTGAACT CTGGTGTGACT CTGGTGTGAACT CTGGTGTGAACT CTGGTGTGAACT CTGGTGTGAACT CTGGTGTGACT CTGCTGTGACT CTGCTGTGACT CTGCTGTGACT CTGCTGTGACT CTGCTGTGACT CTGCTGTGACT CTGCTGTGACT CTTCCCAGTC	51 AGTTGCAAGC TAGGCGAGCC GAAGTATATT TGTTACAGTT ACTTGTGCTA ACACTTGTTTA ACGCTTGTTT ACGCTTGTTT ACGCTAGTATA ACACTTGTTC AAGTAACACT GAACAGCA ACACAGCA ACACAGCA ACACTGACT TCATGGATTAC TCATGGATTAC TCATGGATTAC TCATGGATTAC TCATGGATTC TGCAATTAC TCATGGATTC TGCAATTAC TCATGAATT CCAGAGCATT ACAGGAAAA GAAGGCACCC ACAGCACTC ACAGCACTC ACAGGCACAC GGGACAGCTC AGAGCACTC AGAGCACC AGAGCACTC AGAGCACC AG	120 180 240 300 420 420 660 660 720 780 840 900 1020 1260 1320 1380 1560 1560 1560 1680 1740
50 55 60 65 70	1	11 TATTGACATC CTGATCTTCA GGAATGAGTT ARAATCCCT AGGAATATCG ATTGTGCC AGGAATATCG ATTTTCCAT CTCCTCAGCC GAACACTGGT CTCTCAGCG AACACTGGT TTGGTGTGT TTGGTGTGTT TTGGTGTTTC CTCTAGCGC GAACACTGTT CTCTTGGCTGT TTGGTGTTAC ATTATCAGT CTCTTAGCAG ATGATCTCA GGATGGCTTA TGACACTTAC GGATGGCTTA TGACACTTAC CGGGACTCAG CTTGGTGGG TTGCCTCAGG TTCCCTCAGG TTCCCTCAGG TCCCCAGTCC	1-2712 (underlin 21 AAAAAAAGCA GAATGGTCTA TGATATTAGGTG TGATATTAGGTG TCAGAAAAA TCAAGGAAAA TCAAGGAAAA TCAAGGAACA CCTCTTAGGA TCGTTAGGA TCGTTATAGA CCTCTTAGGA TCGCTGGTATA AGCAGCAATT TCAGGAATGGTG CAGAATGGTG CAGAATGGTG CAGAATGTTG CAGAATGTTG CAGAATGTTG CAGAATGTTGAG AAAGACTA ACAGACTA ACAGACTA ACAGACTA ACAGACTA ACAGACTA TAGTGGTTA TCTGGGATGT AATCATTGTT AGTATCAGTA TCTGGGATG TCTTGGGATG TCTTGGGATG TCTTGGGATG TCTTGGGATG TCTTGGGATG TCTTGGGATG TCTTGGGATG TCTTCGGATG GCACAAGATT	31 AGTGAATTAC AACAAATGTG GAAGATGAGC CTTCTGGCTT ACTGTGGCAA TCTCTTGAAG TTGGAGATTC AGCTTGACAG AATGAGATTC AGCTTGACAG AATGAGATTC AAACAAACG ATGAAGCAAAGG ATGATGAAAGG ATTGTTCAGAA ACTTTCAGAT ACTGAGATC ACTGAGATG ACTGAGATG ACTTTAGTTG CACGACAC ATTAAGTACT TTGGCTTCTG CCACCTAGAA AAAATGATTA TATTCCAAAA AAAATTATTA	41 CAGTCAGTGA AAGTTAGTCA CACTGAGGACAC CACTGCGGA TACTTATCGT AATTGAGTAA CACTTGCCGG CTGCTGACTT GTGAGACAAC TTGCATCAGAG GTGCTTGTCAT CAGAAGAGGC CCTTTTACTC TCATAGTGA ATAAAACTGG GTCTGCATG TGTGCATGT GTGCATGT TGTGCATGAT ACTAGATACCC ACAGGACAG GTACTACATA ACCACAAGA GTCCTGAAT CTTGCTGTAA CTTGCTGTAA CTTGCTGTAA CTTGCTGTAA CTTGCTGAAT CTTGCAATAC CTTGCTGAACT CTTGCTGAACT CTTGCTGAACT CTTGCTGAACT CTTGCTGAAA CTTGGTGTAA AGTCCTAACA AGTCCTAACA AGTCCTAACA AGTCCTACAA	51 AGTTGCAAGC TAGGCGAGCC GAAGTATATT CAGTGTTTA ACTTGTCCA ACTTGTCT ACGCTTGTTT ACGCTTGTTT ACGCTTGTTT ACGCTTGTTT ACGCTAGACAC CTTTGTACA ACACAGGA ACCAAAAACC CTTTGGTATA AATGTTTACT TCATAGGTTCC TAGACAGGTACC TCATGGATTC TCATGGATTC TCCATGAATT TCCAAGAGATT CCAGAGAATT CCAGAGAATT CCAGAGCATT ACACGAGACTT ACACGAGACTT ACACGAGACT ACACGAGACT ACACGAGACT ACACGAGACT ACACGAGACT ACACGAGACT ACACGAGACT ACACGAGACC GGGACACCT ACACGAGACT ACACGAGACT ACACGAGACT ACACGAGACT ACACGAGACT ACACGAGACC GGGACACCT ACACGAGACC GGGACACCT ACACGAGACC ACTCCCACGACAC ACACGAGACC ACACGACACC ACACACC ACACACACC ACACACC ACACACC ACACACC ACACACC ACACACC ACACACC ACACACC ACACACC ACACACC ACACACAC	120 180 240 300 480 540 600 660 780 840 900 1020 1180 1260 1380 1440 1500 1560 1680

5 10	CTAGTGGATMATAACATTIAAAATAACATTIAGATTATTAAAAATAACATTIAGACCTACGAACTCTGCCCTAGTTATTTGACCTAGTTATTTGACCGAACTCTGCCAACTCTTGTACCGAACTCTGCCAACTCTGCCAACTCTGCCAACTCTGCCAACTCTGCCAACTCTGCCAACTCTGCCAACTCTGCAACTCTGCAACTCGAACTCTGCAACTCGAACTCGAACTCGAACTCGAACTCGAACTCGAACTCGAACTCGAACTCGAACACTAGTAACTCGAACACTAGTAACTCGAACACTAGTAACTCGAACACTAGTAACTCGAACACTAGTAACTCGAACACTAGTAACTCGAACACTAGTAACTCGAACACTAGTAACTCGAACACTAGTAACTCGAACACTAGTAACTCGAACACTAGTAACTCGAACACTAGTAACTCGAACACTAGTAACTCGAACACTAGTAACTCGAACACTAGTAACACTAGAACAACAACAACAACAACAACAACAACAACAACAACAA	ATGATTTTCI A AAAATTTCG A CAAAATTTCG C CTACATTAAA A TTATGGATGC C TTATTCATCAT A CAATGTGAT A TGTTCAATGC A GTAATAGAA A TATTCACTC T TCTTTTTGGC A GGAGCAGGGG	A AACCATAATC TAGATTOCAC TAGATTTCCAC GAACTTTCCAC ACCCCAGCT CTCGCAACTGC AATAATCATT TACACCTCGG ACTAAGTTCC ACTTAGGTAC GCTTCAGAAC TCTCACACCTCGG	TCTGCAATCG CTGAGCACGA CAGAGCCTTG AAAGACAGCA GTTTGTGGGA AGACCACAACAA AGATCCCAGA GCAGTTCTTG GCAGTTCTTAGA TCAGTGTGCA	AAGAGGGTAI GTATAGCAGC ATGCCATGCI GAGTAGAACC TTTTGACTAI CTTTGTTTGI TGACCTTCAC ACCAAGTCTGI GATCCATCAI CTGAGAGCCC TAGTGGCAG	AGACATGATC AGGGATTAT ATTAACTITA ATTAACTTTA AGATTTTGTGG AAACTTGATA AAACTTGATA CTTCTGGCGT ATGCTTTGTG GTTTGAGATT GGGACAATTA AAGCATACTG AATTATAAAG ATCATCTTTT	1980 2040 2100 2160 2220 2280 2340 2400 2520 2580 2580 2700
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65 70	CTCGGGATA CCATCAGCC TCTTGTGTCI ACAAATACT TCTCTCACAI ATTAGTATC	TTCTAGAACT GTTATCTGAI TGCTTCCCAT ARGACTGCAG ATACCTCAGI TTCAACTCGI PAV4 Variant 1 Pr	CARTGGTGTG ACTGTCTGAR TGGTGCTGTG CCATGGGCAG GTCTCATGTT AGTAA	CTCTGTGCAA GAACCAAGGA GTGATGGTTT GAAATGTTCT CAGCAGACAA	CTCCCTCAT CACACTCCGA TTGGATTCGT ACTGCTTTCC	TAAGATTATG CATGGCTATT TGACAATTTC	1080 1140 1200
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	CTCGGGATAG CCATCAGCC TCTTGTGTCI ACAAATACT TCTCTCACAA ATTAGTATC SEQ ID NO:2101 Protein Accession 1	TTCTAGAACT GTTATCTGAA TGCTTCCCAT AAGACTGCAA ATACCTCAGI TTCAACTCGI TTCAACTCGI A#: 11 PQUGLPYSM YLLLSVLQFL	C CAATGGTGTG A ACTGTCTGAA CTGTGCTGTGTGTGTGTGTGTGTGTGTGTGTGGGCAG GTCTCATGTTA GTAA Olein sequence; none for 21 KQAGFPLGIL YPFIAMISYN	CTCTGTGCAA GAACCAAGGA GGAACGTTT GGAATGTTCT CAGCAGACAA Ind 31 LLFWVSYVTD LIAGDTLSKV	CTCCCCTCAT CACACTTCCGF TTGGATTCC CACAACTTTC 41 FSLVLLIKGG FQRIPGVDPE	TAAGATTATG CATGGCTATT TAGCAATTTC TACTTTAAAT 51 ALSGTDTYQS NVFIGRHFII	1080 1140 1200 1260
70	CTCGGGATAC CCATCAGCC TCTTGTGTC ACAAATACT TCTCTCACAI ATTAGTATC SEQ ID NO:210 Protein Accession 1 MGYQRQEPVI LVNKTFGFPG GLSTVTFTLP	TTCTAGAACT TGTTATCTGAA TGCTTCCCAT AAGACTGCAA ATACCTCAGA TTCAACTCGA PAV4 Variant 1 Pr 1 # 11 PPQRGLPYSM YLLLSVLQFL LSLYRNIAKL	C CAATGGTGTG A ACTGTCTGAA TGGTGCTGTG C CCATGGGCAG A GTCTCATGTI A GTAA Diein sequence:	CTCTGTGCAA GAACCAAGGA GGTGATGGTTT GGAATGTTCT CAGCAGACAA JI LLPWVSYVTD LLAGDTLSKV TTLILGIVMA	CTCCCTCAT CACACTCCG TTGGATTCG ACTGCTTTCC CACAACTTTC 41 FSLVLLIKGG FSLVLLIKGG FQRIPGVDFE RAISLGPHIF	TAAGATTATG CATGGCTATT TAGCAATTTC TACTTTAAAT 51 ALSGTDTYQS NVFIGRHFII KTEDAWVFAK	1080 1140 1200 1260 60 120 180
70	CTCGGGATAGCCCATCAGCCCTCTTGTGTCTCACAAATACTCTCTCACAAATAGTATCCTCTCACAAATAGTATCCTCTCACAAATAGTATCCTCTCACAAATAGTATCCTCTCACAAATAGTATCCTCACACAAATAGTATCCTCACACACA	TTCTAGAACT TGTTATCTGAA TGCTTATCTGAA ATGCTTCCCAT AAGACTGCAA ATACCTCAGA TTCAACTCGA TTCAACTCGA PAV4 Variant 1 Pr 1 #: 11	CAATGGTGTG ACTGTCTGAA TGGTGCTGTGCCCCCCCCCC	CTCTGTGCAA GAACCAAGGA GGAACGATGTTCT GGAATGTTCT CAGCAGACAA JI LLFWVSYVTD LLAGDTLSKV TTLLLGIVMA TVAKWSRLIH	CTCCCCTCAT CACACTTCCG TTGGATTCCG ACTGCTTTCC CACAACTTTC 41 FSLVLLIKGG FQRIPGVDFE RAISLGPHIP MSIVISVFIC	TAAGATTATG CATAGCTATT TAGATTTC TACTTTAAAT 51 ALSGTDTYQS NVFIGRHFII KTEDANVFAK IFFATCGYLT	1080 1140 1200 1260 60 120 180 240
70	CTCGGGATAGCCCATCAGCCCTCTTGTGTCCTCACAAATACTCTCTCACAAATACTCTCTCACAAATAGTATCCTCTCACAAATAGTATCCTCTCACAAATAGTATCCTCTCACAAATAGTATCCTCACACAAATAGTATCCTCACACACA	TTCTAGAACT TGTTATCTGAA TGCTTATCTGAA ATGCTTCCAGA ATACCTCAGA TTCAACTCGA TTCAACTCGA TTCAACTCGA PAV4 Variant 1 Pr T#: 11 PPQRGLFYSM YLLLSVLOFL LSLYRNIAKL SFAFICHHNS ENYCRNDDLV	C CAATGGTGTG A ACTGTCTGAA A CTGTCTGTGTGTGTGTGTGTGTGTGTGTGTGTGTGTGTG	CTCTGTGCAA GAACCAAGGA GGAACGATCTTCT GGAATGTTCT CAGCAGACAA 31 LLFWVSYVTD LIAGDTLSKV TTLILGIVMA TVAKWSRLIH VILTYPMECF	CTCCCCTCAT CACACTTCGATTCGATTCGATTCGATTCG	TAAGATTATG CATAGCTATT TAGATTTC TACTTTAAAT S1 ALSGTDTYQS NVFIGRHFII KTEDAWFAK IFFARCGYLT FGGNLSSVFH	1080 1140 1200 1260 60 120 180 240 300
70	CTCGGGATAC CCATCAGCC TCTTGTCTC ACAAATACT TCTCTCACAA ATTAGTATC SEQ ID NO:2101 Prolein Accession L MGYQRQEPVI LVMKTFGFPG GLSTVTFTLP PMAIQAVGVM FTGFTQGDLF IVVTVMVITV	TTCTAGAACT TGTTATCTGAA A TGCTTCCCAT AAGACTGCAA ATACCTCAG TTCAACTCGA TTCAACTCGA 11 PPQRGLFYSM YLLLSVLQFL LSLYRNIAKL SFAFICHHIS ENYCRNEDLV ATLVSLLIDC	CAATGGTGTG ACTGTCTGAA TGGTGCTGTGCCCCCCCCCC	CTCTGTGCAA GAACCAAGA GTGATGGTTT GAAATGTTCT CAGCAGACAA 11 LLFWVSYVTD LLFWVSYVTD TLILGIVMA TVAKWSRLIH TVAKWSRLIH VULTYPMECF LCATPLIFII	41 CACACTTTY 41 FSLVLLIKGG FORIPGVDFE RAISLGPHIP MSIVISVFIC VTREVIANVF PSACYLKLSE	TAAGATTATG CATGGCTATT TAGATTTC TACTTTAAAT S1 ALSGTDTYQS NVFIGRHFII KTEDAWFAK IFFATCGYLT FGGNLSSVFH EPRTHSDKIM	1080 1140 1200 1260 60 120 180 240

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_				SEQ ID N	0:211 PAV4 VAR	IANT 2 DNA SEQU	ENCE
5	Nucleic Acid Acco Coding sequence		N62096 1-1203		nces correspond to	o start and stop cod	ions)
	1	11	21	31	41	51	
10	ī	ī	ī	Ĭ	i ·	ĭ	
	ATGGGCTACC	AGAGGCAGGA	GCCTGTCATC	CCGCCGCAGT	TTTCCCTTGT	TTTATTGATA	60
				TACCAGTCTT			120
				CAGTTTTTGT			180
15				AGCAAAGTTT			240
13				TTCATTATTG GCAAAGCTTG			300 360
				GTAATGGCAA			420
				TTTGCAAAGC			480
				CATAACTCCT			540
20				CTTATCCATA			600
				TACTTGACAT			660
				GACCTGGTAA			720
				GAATGCTTTG			780
25				GTTTTCCACA			840
23				ATTGATTGCC TTTATCATTC			900 960
				AAGATTATGT			1020
				ATGGCTATTA			1080
•				GACAATTTCT			1140
30		AGCAGACAAC	ACAACTTTCT	ACTTTAAATA	TTAGTATCTT	TCAACTCGAG	1200
	TAA						
35	SEQ ID NO:212 Protein Accession	PAV4 Variant 2 Pro n #:	otein sequence: none fo	und			
	_				44		
	1	11	21	31	1	51	
40	MCAUDURDAL	PROFESIALIT	PCCALCCTOT	YQSLVNKTFG	•	OFT.VPFTAMT	60
				FIIGLSTVTF			120
				FAKPNAIQAV			180
	EEPTVAKWSR	LIHMSIVISV	FICIFFATCG	YLTFTGFTQG	DLFENYCRND	DLVTFGRFCY	240
15				VFHIVVTVMV			300
45				Kimscvmlpi S tlnisifqle		MAITNTQDCT	360
	Nucleic Acid Acc	ession #-	N62096		0:213 PAV4 VAR	IANT 3 DNA SEQU	ENCE
50	Coding sequence				nces correspond t	o start and stop cod	lons)
	1	11	21	31	41	51	
	i	î.	î	1	i	ĭ	
55	ATGGGCTACC	AGAGGCAGGA	GCCTGTCATC	CCGCCGCAGG	TCAATAAAAC	TTTCGGCTTT	60
				TTTTTGTATC			120
				AAAGTTTTTC			180
				ATTATTGGAC			240
60				AAGCTTGGAA			300 360
UU				ATGGCAAGGG GCAAAGCCCA			420
				AACTCCTTCT			480
	~~~~~~			ATCCATATGT			540
				TTGACATTTA			600
65				CTGGTAACAT			660
	GTCACTGTCA	TTTTGACATA	CCCTATGGAA	TGCTTTGTGA	CAAGAGAGGT	AATTGCCAAT	720
				TTCCACATTG			780
				GATTGCCTCG			840
70				ATCATTCCAT			900
70				ATTATGTCTT GCTATTACAA			960 1020
				AATTTCTCTC			1020
				TTAAATATTA			1000
75				•			
75							
	SEQ ID NO:214 I Protein Accession		oteln seguence: none fo	und			
	1	11	21	31	41	51	
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	•	-	•	•	20	0	
					38	ブ	

5	PENVFIGRHF IPKTEDAWVF ICIFFATCGY VFFGGNLSSV	PPQVNKTFGF IIGLSTVTFT AKPNAIQAVG LTFTGFTQGD FHIVVTVMVI IMSCVMLPIG LNISIFQLE	LPLSLYRNIA VMSFAFICHH LFENYCRNDD TVATLVSLLI	KLGKVSLIST NSFLVYSSLE LVTFGRFCYG DCLGIVLELN	GLTTLILGIV EPTVAKWSRL VTVILTYPME GVLCATPLIF	MARAISLGPH IHMSIVISVF CFVTREVIAN IIPSACYLKL	60 120 180 240 300 360
10				SEQ ID N	0:215 PAV4 VAR	ANT 4 DNA SEQUE	NCE:
	Nucleic Acid Acc Coding sequence		N62096 1-1389	1		o start and stop code	
15	1	11	21	31	41	51	
	ACCCTTGTT	r ctgaacatg	GTATAAAGAG	AAAACCTGT	AGTCTGCTGC	TGACAGAGAA TCTTTTTAAT GAAGCAAGCT	. 60 120 180
20	GGGTTTCCT GTTTTATTG AAAACTTTC ATAGCAATG	T TGGGAATATT A TAAAAGGAGG G GCTTTCCAGG A TAAGTTACAA	GCTTTTATION GGCCCTCTCTCT GGTATCTGCTCT ATATATAGCT	TGGGTTTCAT GGAACAGATA CTCTCTGTTC GGAGATACTT	F ATGTTACAGI A CCTACCAGTC C TTCAGTTTTT F TGAGCAAAGT	CTTTTCCCTT TTTGGTCAAT GTATCCTTTT TTTTCAAAGA	240 300 360 420
25	ACAGTTACC TCCCTCATC TCACTGGGTV ATTCAAGCG	TTACTCTGCC TCTACAGGTTTC CACACATACC TCGGGGTTAT	TTTATCCTTC AACAACTCTC AAAAACAGAA GTCTTTTGC	F TACCGAAATA F ATTCTTGGAA GACGCTTGGC TTTATTTGCC	A TAGCAAAGCT A TTGTAATGGC F TATTTGCAAA C ACCATAACTC	T TGGACTTTCC T TGGAAAGGTC AAGGGCAATT A GCCCAATGCC CTTCTTAGTT A TATGTCCATC	480 540 600 660 720 780
30	GTGATTTCTY TTCACCCAA( AGATTTTGTY GAGGTAATTY	G TATTTATCTO G GGGACTTATT T ATGGTGTCAC G CCAATGTGTT	TATATTCTITE TGAAAATTAC TGTCATTTTC TTTTGGTGGC	GCTACATGTO TGCAGAAATO ACATACCCTI AATCTTTCAT	GATACTTGAC GATGACCTGGT ATGGAATGCTT CGGTTTTCC	ATTTACTGGC ACATTTGGA TGTGACAAGA CATTGTTGTA CCTCGGGATA	840 900 960 1020 1080
35	GTTCTAGAAG TGTTATCTGA ATGCTTCCCA CAAGACTGCA	C TCAATGGTGT A AACTGTCTGA A TTGGTGCTGT A CCCATGGGCA	GCTCTGTGCA AGAACCAAGG GGTGATGGTT AGGAAATGTTC	A ACTCCCTCA S ACACACTCCC TTTTGGATTCC TACTGCTTTC	A TTTTTATCAT F ATAAGATTAT F TCATGGCTAT CTGACAATTT	T TCCATCAGCC T GTCTTGTGTC T TACAAATACT T CTCTCTCACA	1140 1200 1260 1320
40	TTTCAATGA	3 AGTCTCATG1	· TCAGCAGACA	ACACAACTT	CTACTTTAA	TATTAGTATC	1380
45	SEQ ID NO:216   Protein Accession	PAV4 Variant 4 Pro n #:	stein sequence; none for	ınd			
	1	11 	21 	31 	41 	51	
50	GFPLGILLLF IAMISYNIIA SLISTGLTTL YSSLEEPTVA	PFQRDLDDRE WVSYVTDFSL GDTLSKVFQR ILGIVMARAI KWSRLIHMSI	VLLIKGGALS IPGVDPENVF SLGPHIPKTE VISVPICIFF	GTDTYQSLVN IGRHFIIGLS DAWVPAKPNA ATCGYLTFTG	KTFGFPGYLL TVTFTLPLSL IQAVGVMSFA FTQGDLFENY	LSVLQFLYPF YRNIAKLGKV FICHHNSFLV CRNDDLVTFG	60 120 180 240 300
55	VLELNGVLCA	TYPMECFVTR TPLIFIIPSA YCFPDNFSLT	CYLKLSEEPR	THSDKIMSCV	MLPIGAVVMV		360 420
	Shortale Salel Second		101 oct		ID NO:217 PAV9	DNA SEQUENCE	
60	Nucleic Acid Acce Coding sequence		NM_017 1-3501 (		nces correspond to	start and stop codo -	ns)
•	1	11 	21 I	31 ·	41	51 !	
65	GAGAAGCCCA AATTTCCTCC ACATGGGGCT	CCTTCGGGGC CCGATGCCTA GGCTCTCTGA TCCGTGCCCC	CGGAGAGCTG CCGAACGGAT GAACCTGGTG	GACTTCACGG CCAGCTGCAG GTGTCAGTGC	GGGCCGGCCG TTTATAGTCT TGGGGGGATC	CAAGCACAGC GGTCACACGC GGGGGGCCCC	60 120 180 240
70	ACAGGAGCCT GCTGTACGGG GCCCCTGGG	CCTGGCTGCA GGATTGTCAC ACCATCAGAT GTGTGGTCCG GGTGGCGCGG	TGGGGGTCTG GGCCAGCACT GAATAGAGAC	CACACGGGCA GGGGGCACCA ACCCTCATCA	TCGGCCGGCA AGGTGGTGGC ACCCCAAGGG	TGTTGGTGTG CATGGGTGTG CTCGTTCCCT	300 360 420 480 540
75	TACTCGGCCT TTCCGCTTGC ATTGACATCC GAGAACGCCA	TCTTCCTGGT GCCTGGAGTC CTGTCCTGCT CCCAGGCTCA CGGAGACCCT	GGACGACGGC CTACATCTCA CCTCCTGATT GCTCCCATGT	ACACACGGCT CAGCAGAAGA GATGGTGATG CTCCTCGTGG	GCCTGGGGG CGGGCGTGGG AGAAGATGTT CTGGCTCAGG	CGAGAACCGC AGGGACTGGA GACGCGAATA GGGAGCTGCG	600 660 720 780 840
80	GGCGAAGCCC	GAGATCGAAT AGAGGATTAT	CAGGCGTTTC	TTTCCCAAAG	GGGACCTTGA	GGTCCTGCAG	900 960

	GGGTCTGAGG	AATTCGAGAC	CATAGTTTTG	AAGGCCCTTG	TGAAGGCCTG	TGGGAGCTCG	1020
			TGAGCTGCGT				1080
			GGGGGACATC				1140
5			TGACCGGCCT				1200 1260
,			CCTTTTGGAC				1320
			TGCGGAGCTC				1380
			CGCGCCGAGG				1440
10			GAGCATGTAT				1500
10			GCAGGCCCCC				1560
			CATGTACTTC				1620 1680
			GGCGTTCAAG				1740
			GGTGAGGGCT				1800
15			CCAGCTGGCC				1860
			GCTGACACAG				1920
			CGCCTTCTTT				1980
			GGAGCCCACA TGTCGGGACG				2040 2100
20			TCCGGGTTGC				2160
			CTGGGGCGCG				2220
	AGCTACCTGC	TGTTCCTGCT	GCTTTTCTCG	CGGGTGCTGC	TCGTGGATTT	CCAGCCGGCG	2280
			GCTGCTCTAT				2340
25			AGGCGGGGC				2400
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			CTGCATCGAC				2580
			GCTGGGGCCC				2640
20			CTTCCTCGGC				2700
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			CGGGCAGATT				2820 2880
			TGCCAACTGG				2940
			GGTCAACTTG				3000
35			TCTCTACTGG				3060
			GCCCCCCCC				3120
			ACCCCGGAGC				3180
			GGAAGCCGAG ACGCGCTAGG				3240 3300
40			GGACTTGGCA				3360
<i>)</i> -			GGAGCGGGAG				3420
			CTCTGCCTTG	CTGCCCCCAG	GTGGGCCGCC	ACCCCCTGAC	3480
		CCCTGAGCCG CCAAAGAC <u>TG</u>		CTGCCCCCAG	GTGGGCCGCC	ACCCCTGAC	3480
45				CTGCCCCCAG	GTGGGCCGCC	ACCCCTGAC	3480
45	CTGCCTGGGT		<u>A</u>	CTGCCCCAG	GTGGGCCGCC	ACCCCTGAC	3480
45	CTGCCTGGGT	CCAAAGAC <u>TG</u> PAV9 Protein sequ	<u>A</u>		GTGGGCCGCC	ACCCCTGAC	3480
45	SEQ ID NO:2181 Protein Accession	CCAAAGAC <u>TG</u> PAV9 Protein sequ n#:	ence:	und			3480
·	SEQ ID NO:2181 Protein Accession 1	CCAAAGAC <u>TG</u> PAV9 Protein sequ	A ence:		GTGGGCCGCC	ACCCCCTGAC	3480
45	SEQ ID NO:2181 Protein Accession 1	PAV9 Protein sequents:	none fo	und 31	<b>41</b>		
·	SEQ ID NO:218 i Protein Accession  1   MEDAFGAAVV	CCAAAGAC <u>TG</u> PAV9 Protein sequents:  11    TVWDSDAHTT	ence:	und 31   DFTGAGRKHS	41   NFLRLSDRTD		60 120
·	SEQ ID NO:2181 Protein Accession  H HEDAFGAAVV TWGFRAPNLV AVRDHQMAST	PAV9 Protein sequents:  11  TVWDSDAHTT VSVLGGSGGP GGTKVVAMGV	none fo	und 31   DFTGAGRKHS RRGLVRAAQS TLINPKGSFP	41     mplrlsdrtd TGAWIVTGGL ARYRWRGDPE	51     PAAVYSLVTR HTGIGRHYGV DGVQFPLDYN	60 120 180
50	SEQ ID NO:218 i Protein Accession  1   MEDAFGAAVV TWOFRAPNLV AVRDHQMAST YSAFFLVDDG	PAV9 Protein sequents:  11  1 TVWDSDAHTT VSVLGGSGGFV GGTKVVAMGV THGCLGGERR	A none for the control of the contro	und  31   DFTGAGRKHS RRGLVRAAQS TLINPKGSFP QQKTGVGGTG	41     NFLRLSDRTD TGAWIVTGGL ARYRWRGDPE IDIFVLLLLI	51     PAAVYSLVTR   HTGIGRHVGV   DGVQFPLDYN   DGDEKMLTRI	60 120 180 240
·	SEQ ID NO:218 i Protein Accession 1   MEDAFGAAVV TWGFRAPNLV AVRDHQMAST YSAFFLVDDG ENATQAQLPC	CCAAAGAC <u>TG</u> PAV9 Protein sect  #:  11  TVWDSDAHTT VSVLGGSGGP CGTKVVAMGV THGCLGGENR LLVAGSGGAA	none for the state of the state	und  31    DFTGAGRKHS RRGLVRAAQS TLINFKGSFP QQKTGVGGTG LAPGSGGARQ	41     NFLRLSDRTD TGAWIVTGGL ARYRWRGDPE IDIPVLLLLI GEARDRIRRP	51     PAAVYSLVTR   HTGIGRHVGV   DGVQFPLDYN   DGDERMLTRI   FFKGDLEVLQ	60 120 180 240 300
50	SEQ ID NO:2181 Protein Accession  1	PAV9 Protein section #:  11  TVWDSDAHTT VSVLGGSGP GGTKVVAHGV THGCLGGENR LLVAGSGGAA ELLTVYSSED	none for the state of the state	und  31	41     NPLRLSDRTD TGAWIVTGGL ARYRWRGDPE IDIPVLLLLI GEARDRIRRF EASAYLDELR	51 	60 120 180 240 300 360
50	SEQ ID NO:2181 Protein Accession  MEDAFGAAVV TWOFRAPHLV AVRDHQMAST YSAFFLVDDG ENATQAQLEC AQUERIMTRK AQSELFRGDI	PAV9 Protein security  11  TVWDSDAHTT VSVLGGSGGP GGTKVVAMGV THGCLGGENR LLVAGSGGAR LLVAGSGGAR LLVAGSGGAR QWRSFHLEAS	none for the control of the control	und  31   DFTGAGRKHS RRGLVRAAQS TLINPKGSFP QQKTGVGGTG LAPGSGGARQ KALVKACGSS EFVRLLISHG	41   NPLRLSDRTD TGAWIVTGGL ARYRWRGDPE IDIFVLLLLI GEARDRIRRI EASAYLDELR LSLGHFLTFM	51     PAAVYSLVTR HTGIGRHYGV DGVQFPLDYN DGDEKMLTRI FPKGDLEVLQ LAVAWNRVDI RLAVAWNRVDI RLAQLYSAAP	60 120 180 240 300
50 55	SEQ ID NO:218 i Protein Accession  1   MEDAFGAAVV TWOFRAPHLV AVRDHQMAST YSAFFLVDDG ENATQAQLPC AQVERIMTRA AQSELFRGDI SNSLIRNLLD	PAV9 Protein sequents:  11  1 TVWDSDAHTT VSVLGGSGGFV THGCLGGENR LLVAGSGGAA ELLITVYSSEA QWRSFHLEAS QWSFHLEAS	none for the state of the state	und  31   DFTGAGRKHS RRGLVRAAQS TLINFKGSFP QQKTGVGGTG LAPGSGGARQ KALVKACGSS EFVRLLISHG RPPDVGHVLR	41   NFLRLSDRTD TGAWIVTGGL ARYRWRGDPE IDIFVLLLLI GEARDRIRRF EASAYLDEL LSLGHFLTFM MLLGKMCAPR	51 	60 120 180 240 300 360 420
50	SEQ ID NO:218 I Protein Accession  1	CCAAAGACTG  PAY9 Protein sector it:  11  TVWDSDAHTT VSVLGGSGGP GGTKVVAMGV THGCLGGENR LLVAGSGGAA ELLTVYSSED QWRSFHLEAS QASHSAGTKA LLSDKATSPL MARLEPDAEE	none for the state of the state	und  31    DFTGAGRKHS RRGLVRAAQS TLINFKGSFP QQKTGVGGTG LAPGSGGARQ KALVKACGSS EFVRLLISHG RPPDVGHVLR WSDLLLWALL FEGMGVDLFG	41  INFLRLSDRTD TGAWIVTGGL ARYRWRGDPE IDIPVLLLI GEARDRIRRF EASAYLDELR LSLGHFLTPM MLLGKMCAPR LINRAQMAMYF ECYRSSEVRA	51   PAAVYSLVTR HTGIGRHVGV DGVQFPLDYN GDDEKMLTRI FPKGDLEVLQ LAVAWNRVDI RLAQLYSAAP YPSGGAWDPH WEMGSNAVSS ARLLLRRCPL	60 120 180 240 300 360 420 480 540
50 55	SEQ ID NO:218 i Protein Accession  1	PAV9 Protein sequents:  11  1 TVWDSDAHTT VSVLGGSGF GGTKVVAMGV THGCLGGENR LLVAGSGGAR LLVAGSGGAR GUMSFHLEAS QASHSAGTKA LLSDKATSFL MARLEPDAEE MQADARAFFA	A none for the property of the	und  31     DFTGAGRKHS RRGLVRAAQS TLINPKGSFP QQKTGVGGTG LAPGSGGARQ KALVKACGSS EFVRLLISHG RPPDVGHVLR WSDLLLWALL FEGMGVDLFG KWWGDMASTT	41   IMPLRESTATE   MPERESTATE   TARWINTEGEL   ARYKWRGDPE   IDIPVELLE   GEARDRIRRF   EASAYLDEER   LSEGHFETPM   MELGKMCAPR   LNRAQMAMYF   ECYRSSEVRA   PIWALVEAFF	51   PAAVYSLVTR HTGIGRHVGV DGVOFPLDYN DGDEKMLTRI FPKGDLEVLQ LAVAWNRVDI RLAQLYSAAP YPSGGAWDPH WEMGSNAVSS ARLLLRRCPL CPPLIYTRLI	60 120 180 240 300 420 480 540 600
50 55	SEQ ID NO:218 i Protein Accession  1   MEDAFGAAVV TWOFRAPHLV AVRDHQMAST YSAFFLVDDG ENATQAQLPC AQVERIMTRK AQSELFRGDI SNSLIRNLLD PGQGFGESMY ALGACLLLRV WGDATCLQLA TFRKSEEEPT	PAV9 Protein sectors:  11  1 VSVLGGSGGP GGTKVVAMGV THGCLGGENR LLVAGSGGAA ELLTVYSSEA ELLTVYSSEA QASHSAGTKA LLSDKATSPL MARLEPDAE MQADARAFFA REELEFDMDS	A none for n	und  31	41   NFLRLSDRTD TGAWIVTGGL ARYRWRGDPE IDIFVLLLLI GEARDRIRRF EASAYLDEL LSLGHFLTPM MLLGKMCAPR LNRAQMAMYF ECYRSSEVRA FIWALVLAFF VFRQSGRPGC	51	60 120 180 240 300 360 420 480 540 600 720
50 55 60	SEQ ID NO:218 I Protein Accession  1   MEDAFGAAVV TWOFRAPNLV AVROHOMAST YSAFFLVDDG ENATOAQLPC AQVERIMTRK AQSELFRGDI SNSLIRNLLD PGQGFGESMY ALGACLLLRV WGDATCLQLL LTFRKSEEEPT LRWFHFWGA	CCAAAGACTG  PAV9 Protein secutive:  11  1 TVWDSDAHTT VSVLGGSGGP CGTKVVAMGV THGCLGGENR LLVAGSGGAA ELLVAGSGGAA ELLVAGSGGAA ELLVAGSGAA ELLVAGSGGAA ELVAGSGGAA	none for the control of the control	und  31	41  INFLRLSDRTD TGAWIVTGGL ARYRWRGDPE IDIPVLLLLI GEARDRIRRP EASAYLDELR LSLGHFLTPM MLLGKMCAPR LNRAQMAMYF ECYRSSEVRA PIWALVLAFP VPRQSGRPGC PPGSLELLLY	51   PAAVYSLVTR HTGIGRHVGV DGVQFPLDYN DGDERMLTRI FFKGDLEVLQ LAVAWNRVDI RLAQLYSAAP YPSGGAMDPH WEMGSNAVSS ARLLLRRCPL CPFLIYTRLI CGGRCGGRRC FWAFTLLCEE	60 120 240 300 360 420 540 600 660 720 780
50 55	SEQ ID NO:218 I Protein Accession  1	CCAAAGACTG  PAV9 Protein sector it:  11  TVWDSDAHTT VSVLGGSGGP GGTKVVAMGV THGCLGGENR LLVAGSGGAA ELLTVYSSED QWRSFHLEAS QASHSAGTKA LLSDKATSPL MARLEPDAEE MQADARAFFA REELEPDMDS PVTIPMGNVV SLASGGPGPG	A none for n	und  31	41  INFLRLSDRTD TGAWIVTGGL ARYRWRGDPE EDIPVLLLI GEARDRIRRF EASAYLDELR LSLGHFLTPM MLLGKMCAPR LINRAQMAMYF ECYRSSEVRA FIWALVLAFF VPRQSGRPGC PPGSLELLLY LVALTCFLLG	51   PAAVYSLVTR HTGIGRHVGV DGVQFPLDYN GDDEKMLTRI FPKGDLEVLQ LAVAWNRVDI RLAQLYSAAP YPSGGAWDPH WEMGSNAVSS ARLLRCPL CPPLIYTRLI CGGRCGGRC FWAFTLLCEE VGCRLTPGLY	60 120 180 240 300 360 420 480 540 600 720
50 55 60	SEQ ID NO:218 i Protein Accession  1   MEDAFGAAVV TWOFRAPHLV AVROHOMAST YSAFFLVDDG ENATQAQLPC AQVERIMTRK AQSELFRGDI SNSLIRNLLD PGQGFGESMY ALGACLLLRV WGDATCLQLA TFRKSEEEPT LRWHHFWGA LRQGLSGGG HLGRTVLCID EGLLRPRDSD	PAV9 Protein sectors:  11  1 VSVLGGSGGP GGTKVVAMGV GTGCGGENR LLVAGSGGAA ELLTVYSSEL QASHSAGTKA LLSDKATSPL MARLEPDAES MARLEPDAES PVTIFMGNVV SLASGGFGPG FWFTVRLH FFSILRRVFY	A none for the control of the contro	31	41  INFLRESDRTD TGAMIVTGGL ARYRWRGDPE IDIFVLLLLI GEARDRIRRF EASAYLDEL LSLGHFLTFM MLLGKMCAPR LNRAQMAMYF ECYRSSEVAR FIWALVLAFF VPRQSGRPGC PPGSLELLLY LVALTCFLLG EHSNCSSEPG	51	60 120 180 240 300 360 420 480 540 600 720 780 840 900 960
50 55 60	SEQ ID NO:2181 Protein Accession  1   MEDAFGAAVV TWOFRAPHLV AVROHOMAST YSAFFLVDDG ENATOAQLPC AQVERIMTRK AQSELFRGDI SNSLIRNLLD PGQGFGESMY ALGACLLLRV WGDATCLQLL LRV WGDATCLQLL LRV HFWFHFWGA LRQSLSGGGG HLGRTVLCID GGLLSGGGG HLGRTVLCID GTCVSQYANW	CCAAAGACTG  PAV9 Protein sectors:  11  1 TVWDSDAHTT VSVLGGSGGP CGTKVVAMGV THGCLGGENR LLVAGSGGAA ELLTVYSSED QWRSFHLEAS QASHSAGTKA LLSDKATSPL MARLEPDAEE MQADARAFFA REELEFDMDS PVTIPMGNVV SLASGFGFGF FMVTVRLLLVIFL LVVLLLVIFL	none for the control of the control	und  31    DFTGAGRKHS RRGLVRAAQS TLINFKGSFP QQKTGVGGTG LAPGSGGARQ KALVKACGSS EFVRLLISHG RPPDVGHVLR WSDLLLWALL FEGMGVDLFG KWWGDMASTI ADPAEKTPLG RVLLVDFQPA YLADSWNQCM KIVIVSKMM PQEDMDVALM LIAMFSYTFG	41  INFLRLSDRTD TGAWIVTGGL ARYRWRGDPE IDIPVLLLLI GEARDRIRRP EASAYLDELR LSLGHFLTPM MLLGKMCAPR LNRAQMAMYF ECYRSSEVRA PIWALVLAFP VPRQSGRPGC PPGSLELLLY LVALTCFLLG DVFFLFFLG KVQGNSDLYW	51   PAAVYSLVTR HTGIGRHVGV DGVQFPLDYN DGDERMLTRI FFKGDLEVLQ LAVAWNRVDI RLAQLYSAAP YPSGGAMDPH WEMGSNAVSS ARLLLRRCPL CPFLIYTRLI CGGRCGGRRC FWAFTLLCEE VGCRLTPGLY VWLVAYGVAT KAQRYRLIRE	60 120 240 300 360 420 540 600 660 720 780 840 900 960 1020
50 55 60	SEQ ID NO:218 I Protein Accession  I	PAV9 Protein secunit:  11  1 TVWDSDAHTT VSVLGGSGGP GGTKVVAMGV THGCLGGENR LLVAGSGGAR LLVAGSGGAR QMRSFHLEAS QASHSAGTKA LLSDKATSFL MQADARAFFA REELEFDMDS PVTIPMGNV SLASGGPGG FMVFTVRLIH FPSILRRVFTY LVVLLLVIFL PIVISHLRLL	A none for the control of the contro	JUND 31   DFTGAGRKHS RRGLVRAAQS TLINPKGSFP QQKTGVGGTG LAPGSGGARQ KALVKACGSS EFVRLLISHG RPPDVGHVLR WSDLLLWALF FEGMGVDLFG KWWGDMASTT ADPAEKTPLG RVLLVDFQPA YLADSWNQCD KIVIVSKMMK PQEDMDVALM LIAMFSYTFG PQFSSPALEH	41    MFLRLSDRTD TGAWIVTGGL ARYRWRGDPE IDIPVLLLI GEARDRIRRF EASAYLDELR LSLGHFLTFM MLLGKMCAPR LNRAQMAMYF ECYRSSEVRA PIWALVLAFF VPRQSGRPGC PPGSLELLLY LVALTCFFLLG DVFFFLFFLG EHSNCSSEPG KVQGNSDLYW FRVYLSKEAE	51  PAAVYSLVTR HTGIGRHVGV DGVQFPLDYN DGDEKMLTRI PFKGDLEVLQ LAVAWNVDI RLAQLYSAAP YPSGGANDPH WEMGSNAVSS ARLLLRRCPL CPPLIYTRLI CGGRCGGRC FWAFTLLCEE VGCRLTFGLY VWLVAYGVAT FWAHPPCAQA KAQRYRLIRE RKLLTWESVH	60 120 180 240 300 360 420 540 600 660 720 780 840 900 960 1020 1080
50 55 60	SEQ ID NO:218 i Protein Accession  1   MEDAFGAAVV TWOFRAPHLV TWOFRAPHLV AVRDHQMAST YSAFFLVDDG ENATOAQLPC AQVERIMTKA AQSELFRGDI SNSLIRNLLD PGQOFGESMY ALGACLLLRV WGDATCLQLA TFRKSEEEPT LRRWFHFWGA LRQGLSGGGG HLGRTVLCID EGLLRPRDSD GTCVSQYANW PHSRPALAPP KEMFLLARAR	PAV9 Protein sequents:  11  1 TVWDSDAHTT VSVLGGSGGP GGTKVVAMGV THGCLGGENR LLVAGSGGAA ELLTVYSSED QWASFHLEAS QASHSAGTKA LLSDKATSPL MARLEPDAEE MQADARAFFA REELEPDMDS PVTIPMGNVV SLASGGPGPG FMVFTVRLH FPSILRRVFY LVVLLLVIFL LVISHLRLL DKRESDSERL	A none for n	JUND 31   DFTGAGRKHS RRGLVRAAQS TLINPKGSFP QQKTGVGGTG LAPGSGGARQ KALVKACGSS EFVRLLISHG RPPDVGHVLR WSDLLLWALF FEGMGVDLFG KWWGDMASTT ADPAEKTPLG RVLLVDFQPA YLADSWNQCD KIVIVSKMMK PQEDMDVALM LIAMFSYTFG PQFSSPALEH	41    MFLRLSDRTD TGAWIVTGGL ARYRWRGDPE IDIPVLLLI GEARDRIRRF EASAYLDELR LSLGHFLTFM MLLGKMCAPR LNRAQMAMYF ECYRSSEVRA PIWALVLAFF VPRQSGRPGC PPGSLELLLY LVALTCFFLLG DVFFFLFFLG EHSNCSSEPG KVQGNSDLYW FRVYLSKEAE	51  PAAVYSLVTR HTGIGRHVGV DGVQFPLDYN DGDEKMLTRI PFKGDLEVLQ LAVAWNVDI RLAQLYSAAP YPSGGANDPH WEMGSNAVSS ARLLLRRCPL CPPLIYTRLI CGGRCGGRC FWAFTLLCEE VGCRLTFGLY VWLVAYGVAT FWAHPPCAQA KAQRYRLIRE RKLLTWESVH	60 120 240 300 360 420 540 600 660 720 780 840 900 960 1020
<ul><li>50</li><li>55</li><li>60</li><li>65</li></ul>	SEQ ID NO:218 i Protein Accession  1   MEDAFGAAVV TWOFRAPHLV TWOFRAPHLV AVRDHQMAST YSAFFLVDDG ENATOAQLPC AQVERIMTKA AQSELFRGDI SNSLIRNLLD PGQOFGESMY ALGACLLLRV WGDATCLQLA TFRKSEEEPT LRRWFHFWGA LRQGLSGGGG HLGRTVLCID EGLLRPRDSD GTCVSQYANW PHSRPALAPP KEMFLLARAR	PAV9 Protein secunit:  11  1 TVWDSDAHTT VSVLGGSGGP GGTKVVAMGV THGCLGGENR LLVAGSGGAR LLVAGSGGAR QMRSFHLEAS QASHSAGTKA LLSDKATSFL MQADARAFFA REELEFDMDS PVTIPMGNV SLASGGPGG FMVFTVRLIH FPSILRRVFTY LVVLLLVIFL PIVISHLRLL	A none for n	und  31    DFTGAGRKHS RRGLVRAAQS TLINFKGSFP QQKTGVGGTG LAPGSGGARQ KALVKACGSS EFVRLLISHG RPPDVGHVLR WSDLLLWALL FEGMGVDLFG KWWGDMASTI ADPAEKTPLG RVLLVDFQPA YLADSWNQCM KIVIVSKMK PQEDMDVALM LIAMFSYTFG PQPSSPALEH LKQLGHIREY	41  INFLRLSDRTD TGAMIVTGGL ARYRWRGDPE IDIPVLLLLI GEARDRIRRP EASAYLDELR LSLGHFLTPM MLIGKMCAPR LNRAQMAMYF ECYRSSEVRA PIWALVLAFP VPRQSGRPGC PPGSLELLLY LVALTCFLLG DVFFLFFLG KVQGNSDLYW FRVYLSKEAE EQRLKVLERE	51   PAAVYSLVTR HTGIGRHVGV DGVQFPLDYN DGDERMLTRI FFKGDLEVLQ LAVAWNRVDI RLAQLYSAAP YPSGGAMDPH WEMGSNAVSS ARLLLRRCPL CPFLIYTRLI CGGRCGGRRC FWAFTLLCEE VGCRLTPGLY VWLVAYGVAT KAQRYRLIRE RKLLTWESVH VQQCSRVLGW	60 120 240 300 360 420 540 6600 6600 720 780 840 900 960 1020 1080 1140
<ul><li>50</li><li>55</li><li>60</li><li>65</li></ul>	SEQ ID NO:218 I Protein Accession  I	PAV9 Protein section :  11  1 TVWDSDAHTT VSVLGGSGFP GGTKVVAMGV THGCLGGENR LLVAGSGGAR LLVAGSGGAR QWRSFHLEAS QASHSAGTKA LLSDKATSFL MARLEPDAEE MQADARAFFA REELEFDMDS PVTIFMGNV SLASGGPGPG FMVFTVRLLH FPSILRRVFY LVVLLLVIFL DKRESDSERL LPPGGPPPPD	A none for none for none for the first state of the	und  31    DFTGAGRKHS RRGLVRAAQS TLINFKGSFP QQKTGVGGTG LAPGSGGARQ KALVKACGSS EFVRLLISHG RPPDVGHVLR WSDLLLWALL FEGMGVDLFG KWWGDMASTI ADPAEKTPLG RVLLVDFQPA YLADSWNQCM KIVIVSKMK PQEDMDVALM LIAMFSYTFG PQPSSPALEH LKQLGHIREY	41  INFLRLSDRTD TGAMIVTGGL ARYRWRGDPE IDIPVLLLLI GEARDRIRRP EASAYLDELR LSLGHFLTPM MLIGKMCAPR LNRAQMAMYF ECYRSSEVRA PIWALVLAFP VPRQSGRPGC PPGSLELLLY LVALTCFLLG DVFFLFFLG KVQGNSDLYW FRVYLSKEAE EQRLKVLERE	51  PAAVYSLVTR HTGIGRHVGV DGVQFPLDYN DGDEKMLTRI PFKGDLEVLQ LAVAWNVDI RLAQLYSAAP YPSGGANDPH WEMGSNAVSS ARLLLRRCPL CPPLIYTRLI CGGRCGGRC FWAFTLLCEE VGCRLTFGLY VWLVAYGVAT FWAHPPCAQA KAQRYRLIRE RKLLTWESVH	60 120 240 300 360 420 540 6600 6600 720 780 840 900 960 1020 1080 1140
<ul><li>50</li><li>55</li><li>60</li><li>65</li></ul>	SEQ ID NO:218 i Protein Accession  1   MEDAFGAAVV TWOFRAPHLY TWOFRAPHLY YSAFFLVDDG ENATOAQLPC AQVERIMTRK AQSELFRGDI SNSLIRNLLD PGQOFGESMY ALGACLLLRV WGDATCLQLA TFRKSEEEPT LRRWFHFWGA LRQGLSGGG HLGRTVLCID EGLLRPRDSD GTCVSQYANW FHSRPALAPP KENFLLARAR VAEALSRSAL	PAV9 Protein sequents:  11  1 TVWDSDAHTT VSVLGGSGGA ELLTVYSSED QWRSFHLEAS QASHSAGTKA LLSDKATSFL LLSDKATSFL MARLEFDAEE MQADARAFFA REELEFDMDS FVTIFMENVV SLASGGPGPG FMVFTVRLLH FPSILRRVFY LVVLLLVIFL DKRESDSERL LPPGGPPPPD	A none for none for none for none for the first state of the first sta	Jand  31	41    NFLRLSDRTD TGAWIVTGGT ARYRWRGDPE IDIFVLLLI GEARDRIRRF EASAYLDELL LIGHMCAPR LNRAQMAMYF ECYRSSEVRA PIWALVLAFF VPRQSGRPGC PPGSLELLLY LVALTCFLLG EHSNCSSEPG KVQGNSDLYM FRVYLSKEAE EQRLKVLERE	51	60 120 240 300 360 420 540 6600 6600 720 780 840 900 960 1020 1080 1140
<ul><li>50</li><li>55</li><li>60</li><li>65</li></ul>	SEQ ID NO:218 I Protein Accession  I	PAV9 Protein sequents:  11  1 TVWDSDAHTT VSVLGGSGGA ELLTVYSSED QWRSFHLEAS QASHSAGTKA LLSDKATSFL LLSDKATSFL MARLEFDAEE MQADARAFFA REELEFDMDS FVTIFMENVV SLASGGPGPG FMVFTVRLLH FPSILRRVFY LVVLLLVIFL DKRESDSERL LPPGGPPPPD	A none for none for none for none for the first state of the first sta	und  31    DFTGAGRKHS RRGLVRAAQS TLINFKGSFP QQKTGVGGTG LAPGSGGARQ KALVKACGSS EFVRLLISHG RPPDVGHVLR WSDLLLWALL FEGMGVDLFG KWWGDMASTI ADPAEKTPLG RVLLVDFQPA YLADSWNQCM KIVIVSKMK PQEDMDVALM LIAMFSYTFG PQPSSPALEH LKQLGHIREY	41    NFLRLSDRTD TGAWIVTGGT ARYRWRGDPE IDIFVLLLI GEARDRIRRF EASAYLDELL LIGHMCAPR LNRAQMAMYF ECYRSSEVRA PIWALVLAFF VPRQSGRPGC PPGSLELLLY LVALTCFLLG EHSNCSSEPG KVQGNSDLYM FRVYLSKEAE EQRLKVLERE	51	60 120 240 300 360 420 540 6600 6600 720 780 840 900 960 1020 1080 1140
<ul><li>50</li><li>55</li><li>60</li><li>65</li><li>70</li></ul>	SEQ ID NO:218 i Protein Accession  1   MEDAFGAAVV TWOFRAPHLY TWOFRAPHLY YSAFFLVDDG ENATOAQLPC AQVERIMTRK AQSELFRGDI SNSLIRNLLD PGQOFGESMY ALGACLLLRV WGDATCLQLA TFRKSEEEPT LRRWFHFWGA LRQGLSGGG HLGRTVLCID EGLLRPRDSD GTCVSQYANW FHSRPALAPP KENFLLARAR VAEALSRSAL	PAV9 Protein sequents:  11  1 TVWDSDAHTT VSVLGGSGGA ELLTVYSSED QWRSFHLEAS QASHSAGTKA LLSDKATSFL LLSDKATSFL MARLEFDAEE MQADARAFFA REELEFDMDS FVTIFMENVV SLASGGPGPG FMVFTVRLLH FPSILRRVFY LVVLLLVIFL DKRESDSERL LPPGGPPPPD	A none for none for none for none for the first state of the first sta	Jand  31	41    NFLRLSDRTD TGAWIVTGGT ARYRWRGDPE IDIFVLLLI GEARDRIRRF EASAYLDELL LIGHMCAPR LNRAQMAMYF ECYRSSEVRA PIWALVLAFF VPRQSGRPGC PPGSLELLLY LVALTCFLLG EHSNCSSEPG KVQGNSDLYM FRVYLSKEAE EQRLKVLERE	51	60 120 240 300 360 420 540 6600 6600 720 780 840 900 960 1020 1080 1140
<ul><li>50</li><li>55</li><li>60</li><li>65</li><li>70</li></ul>	SEQ ID NO:218 i Protein Accession  1   MEDAFGAAVV TWOFRAPHLY AVRDHQMAST YSAFFLVDDG ENATOAQLPC AQVERIMTKA AQSELFRGDI SNSLIRNLLD PGQGFGESMY ALGACLLLRV WGDATCLQLA TFRKSEEEPT LRRWFHFWGA LRQGLSGGGG HLGRTVLCID EGLLRPRDSD GTCVSQYANW PHSRPALAPP KEMFLLARAR VAEALSRSAL  Nucleic Acid Acce Coding sequence	PAV9 Protein sequents:  11  1 TVWDSDAHTT VSVLGGSGGAA ELLTVYSSED CWRSFHLEAS QASHSAGTKA LLSDKATSPL MARLEPDAEE MQADARAFFA REELEPDMDS PVTIPMGNVV SLASGGPPG FMVFTVRLLH FPSILRRVFY LVVLLLVIFL LVVISHLFLL DKRESDSERL LPPGGPPPPD  ESSION #:	A none for none for none for none for the first state of the first sta	and  31	41    NFLRLSDRTD TGAWIVTGGI ARYRWRGDPE IDI FVLLLLI GEARDRIRRF EASAYLDELL LSLGHFLTPM MLLGKMCAPR LNRAQMAMYF ECYRSSEVRA FIWALVLAFF VPRQSGRPGC PPGSLELLLY LVALTCFLLG DVFFFLFFLG EHSNCSSEPG KVQGNSDLYW FRVYLSKEAE EQRLKVLERE    ID NO:219 PBF1 respond to start at 41	51   PAAVYSLVTR HTGIGRHVOV DGVQFPLDYN DGDEKMLTRI FFKGDLEVLQ LAVAWNRVDI RLAQLYSAAP YPSGGAWDPH WEMGSNAVSS ARLLLRRCPL CPPLIYTRLI CGGRCGGRRC FWAFTLLCEE VGCRLTPGLY VWLVAYGVAT FWAHPPGAQA KAQRYRLIRE RKLLTWESVH VQQCSRVLGW  DNA SEQUENCE dd stop codons) 51	60 120 180 240 300 360 420 480 540 660 720 780 840 900 960 1020 1080
<ul><li>50</li><li>55</li><li>60</li><li>65</li><li>70</li></ul>	SEQ ID NO:218 i Protein Accession  1   MEDAFGAAVV TWOFRAPNILV AVROHOMAST YSAFFIVDDG ENATOAQLPC AQVERIMTRK AQSELFRGDI SNSLIRNLLD PGQGFGESMY ALGACLLLRV WGDATCLQLA TFRKSEEEPT LRWPHPWGA LRQGLSGGGG HLGRTVLGID EGLLRPDSD GTCVSQYANW PHSRPALAPR VAEALSRSAL  Nucleic Acid Acce Coding sequence  1   ATGGAGCCCC	PAV9 Protein sectors:  11  1 TVWDSDAHTT VSVLGGSGGP GGTKVVAMGV GGTKVVAMGV THGCLGGENR LLVAGSGGAA ELLTVYSSED QWRSFHLEAS QWRSFHLEAS QWRSFHLEAS PAVE MARLEPDAEF LUSHLRRVFY LVVLLLVLIFL LVVLLLVIFL LVVLLLVIFL LPPGGPPPPD  SSSION #:  11  GGGGCGCCTCGT	A none for the part of the par	and  31	41  INFLRLSDRTD TGAMIVTGGL ARYRWRGDPE IDIFVLLLI GEARDRIRRF EASAYLDELR MILGKMCAPR LNRAQMAMYF ECYRSSEVRA PIWALVLAFF VPRQSGRPGC PPGSLELLLY LVALTCFLLG EHSNCSSEPG KVQGNSDLYW FRVYLSKEAE EQRLKVLERE  DID NO:219 PBF1 respond to start at 41  I TCAGCCTGTG	51   PAAVYSLVTR HTGIGRHYGV DGVQPPLDYN DGDEXMLTRI FPKGDLEVLQ LAVAWNRVDI RLAQLYSAAP YPSGGAWDPH WEMGSNAVSS ARLLERCPL CPPLIYTRLI CGGRCGGRRC FWAFTLLCEE VGCRLTFGLY VWLVAYGVAT FWAHPPGAQA KAQRYRLIRE RKLLTWESVH VQQCSRVLGW  DNA SEQUENCE ad slop codons) 51   CTCCCTGGGG	60 120 180 240 300 360 420 480 540 660 720 780 840 960 1020 1080 1140
<ul><li>50</li><li>55</li><li>60</li><li>65</li><li>70</li><li>75</li></ul>	SEQ ID NO:2181 Protein Accession  1   MEDAFGAAVV TWOFRAPRILV AVROHOMAST YSAFFLVDDG ENATOAQLPC AQVERIMTRK AQSELFRGDI SNSLIRNLLD PGQGFGESMY ALGACLLLRV WGDATCLQLA TFRKSEEEPT LRWFHFWGA LRQGLSGGGG HLGRTVLCID GTCVSQYANW FHSRPALAPP KENFLLARA VAEALSRSAL  Nucleic Acid Acce Coding sequence  1   ATGGAGCCGC CTGCTCGTCA	PAV9 Protein sectors:  11  1 VSVIGGSGGP GGTKVVAMGV THGCLGGENR LLVAGSGGAA ELLITVYSSED QWRSFHLEAS QWRSFHLEAS QWRSFHLEAS PAVIFMGNVV SLASGGFGFR ELEFDMDS FVTIFMGNVV SLASGGFGFR LVVLLLVIFL FVSLIRRVFY LVVLLLVIFL FVSLIRRVFY LVVLLLVIFL FSSION #:  11  1  1  1  1  1  1  1  1  1  1  1	A none for the property of the	und  31	41  INFLRLSDRTD TGAMIVTGGL ARYRWRGDPE IDIPVLLLI GEARDRIRRP EASAYLDELR LSLGHFLTPM MILGKMCAPR LNRAQMAMYF ECYRSSEVRA PIWALVLAFP VPRQSGRPGC PPGSLELLLY LVALTCFLLG EHSNCSSEPG KVQGNSDLYW FRVYLSKEAE EQRLKVLERE A ID NO.219 PBF1 TESPOND to start at  41  I TCAGCCTGTG CCGACCCCCG	51   PAAVYSLVTR HTGIGRHVGV DGVQPPLDYN DGDERMLTRI FPKGDLEVLQ LAVAWNRVDI RLAQLYSAAP YPSGGAMDPH WEMGSNAVSS ARLLLRRCPL CPPLIYTRLI CGGRCGGRRC FWAFTLLCEE VGCRLTFGLY VWLVAYGVAT FWAHPPGAQA KAQRYRLIRE RKLLTWESVH VQQCSRVLGW  DNA SEQUENCE ad stop codons) 51   CTCCCTGGGG GCGCCACAAG	60 120 240 300 360 420 540 600 660 720 780 840 900 960 1020 1080 1140
<ul><li>50</li><li>55</li><li>60</li><li>65</li><li>70</li></ul>	SEQ ID NO:2181 Protein Accession  1   MEDAFGAAVV TWOFRAPRILV AVROHOMAST YSAFFLVDDG ENATOAQLPC AQVERIMTRK AQSELFRGDI SNSLIRNLLD PGQGFGESMY ALGACLLLRV WGDATCLQLA TFRKSEEEPT LRWFHFWGA LRQGLSGGGG HLGRTVLCID GTCVSQYANW FHSRPALAPP KENFLLARA VAEALSRSAL  Nucleic Acid Acce Coding sequence  1   ATGGAGCCGC CTGCTCGTCA	PAV9 Protein sectors:  11  1 VSVIGGSGGP GGTKVVAMGV THGCLGGENR LLVAGSGGAA ELLITVYSSED QWRSFHLEAS QWRSFHLEAS QWRSFHLEAS PAVIFMGNVV SLASGGFGFR ELEFDMDS FVTIFMGNVV SLASGGFGFR LVVLLLVIFL FVSLIRRVFY LVVLLLVIFL FVSLIRRVFY LVVLLLVIFL FSSION #:  11  1  1  1  1  1  1  1  1  1  1  1	A none for the part of the par	und  31	41  INFLRLSDRTD TGAMIVTGGL ARYRWRGDPE IDIPVLLLI GEARDRIRRP EASAYLDELR LSLGHFLTPM MILGKMCAPR LNRAQMAMYF ECYRSSEVRA PIWALVLAFP VPRQSGRPGC PPGSLELLLY LVALTCFLLG EHSNCSSEPG KVQGNSDLYW FRVYLSKEAE EQRLKVLERE A ID NO.219 PBF1 TESPOND to start at  41  I TCAGCCTGTG CCGACCCCCG	51   PAAVYSLVTR HTGIGRHYGV DGVQPPLDYN DGDEKMLTRI FPKGDLEVLQ LAVAWNRVDI RLAQLYSAAP YPSGGANDPH WEMGSNAVSS ARLLERCPL CPPLIYTRLI CGGRCGGRRC FWAFTLLCEE VGCRLTPGLY VWLVAYGVAT FWAHPPGAQA KAQYRLIRE RKLLTWESVH DNA SEQUENCE ad slop codons) 51   CTCCCTGGGG GCGCCACAAG CCGCCTGATG	60 120 180 240 300 360 420 480 540 660 720 780 840 960 1020 1080 1140

5	GGCCCGGGC GCCGAGTGCG CTGGGCATGC GCCATCAAGT AAGACCATAC CTCGGCATGG TGGGAGGAGA TGCACCATTT AAGCTAATTT GCCTGGTGCA	ACCTGCCGCT GCGCCGACCC GCCGGCCCCT ACCGGGACAT ACCACTTTTC AGCAAGATGA CCGTAGCCGT GCTTGACCCA CCTCTGTAC ATAGCCTGCC GTTTAGCCTTGCCT CCAAGATTGC	CGAGTCCTGG CTTCGCCACC CGACACCCTC TCAGCACCTG CTTGCCACCTG CCTTCTCTGC GCACGTGGCT TTATGCCGCC TGCTGATGTG TATTGTGGCA	CGCTCGCTCC TACTCGGGCC ATCCTGAAAG CGCTTGCGAA CTTCATTTAA GGCTGCATTG GGACTCCTGT GAACATGGTT GCTGGAGGTC	TGGGGCTCGG TCTGGAGGAA GTATTGCGCA ACATTCCTTT GAAGAATCAC TGGCCACAGT TCCTCATGAC ATGATTGAA ACAGCTGGTC TCTGCATCGC	CGGGCTGGAC GTGCTACTTC GCGATGCACG TAATTTAACC TGCTGGCTTC CAGTTTCTTC AGGGATATTT CCGGCTCCCA CATCTTTTGC TTATCCGTTT	240 300 360 420 480 540 600 660 720 780 840
15	SEQ ID NO:220   Protein Accession	PBF1 Protein sequ n#:	none found				
20 25	PLSHLPLRDS LGIDRDIDTL LGMAVAVLLC	11   SLGLSLCSLG PPLGRRLLPG ILKGIAQRCT GCIVATVSFF EHGYSWSIFC	GPGRADPESW AIKYHFSQPI WEESLTQHVA	RSLLGLGGLD RLRNIPFNLT GLLFLMTGIF	AECGRPLFAT KTIQQDEWHL CTISLCTYAA	YSGLWRKCYF LHLRRITAGF SISYDLNRLP	60 120 180 240
				SE	Q ID NO:221 PCI4	DNA SEQUENCE	
	Nucleic Acid Acc		NM_016570				
30	Coding sequence	ĸ	1- 1134 (under	fined sequences o	отеѕропо то ѕтал	and stop codons)	
	1	11	21	31	41	51	
35	AAGGTTCCTG TTTACAACTA AAGTATGAAT ACTGTTGCCA	TGAATCGGAA AGAGCTATGT TGGCTTTATT ACGAAGTAGA TGAAGTGTCA	AGAGACTTCA AACCATAATG CAAGGATTTT ATATGTTGGA	GCCAGTGGAG GAATTCTCAG TCTAGCAAAT GCGGATGTAT	GTACAGTTTC TATATCAAGA TAAGAATTAA TGGATTTAGC	TCTAATAGCA TACATGGATG TATAGATATT AGAAACAATG	60 120 180 240 300
40	AAAGAGTGGC CAAGATGTGA GATGATTCAT	CAGATGGTTT AGAGGATGCT TATTTAAAAG CACAGTCTCC ATTTTCACAT	GCAGCTGATT TGCTTTTAAA AAATGCATGC	CAGAGTAGGC AGTACATCAA AGAATTCATG	TACAAGAAGA CAGCTCTTCC GCCATCTATA	GCATTCACTT ACCAAGAGAA TGTCAATAAA	360 420 480 540 600
45	CATTTGCAG TCTTTTGGAG ATAGATCACA TATAAAATAT CATGCTGCAG	CACTTGTCAA AGCTTGTTCC ACCAGATGTT CAGCAGACAC GCAGCCATGG	CCATGAATCT AGCAATATTT CCATCAGTTT AGTCTCTGGG	TACAATTTTT AATCCTTTAG ATTACAGTTG TCTGTGACAG ATATTTATGA	CTCATAGAAT ATGGAACTGA TGCCAACAAA AAAGGGAACG AATATGATCT	AGATCATTTG AAAAATTGCT ACTACATACA TATCATTAAC CAGTTCTCTT	660 720 780 840 900 960
50	ATTGTTGGAG GAAATAATTT	TTACTGAGGA GAATCTTTTC GCTGTCGTTT ACACAGACAA	AACAACAGGC CAGACTTGGA	ATGTTACATG TCCTATAAAC	GAATTGGAAA CTGTCAATTC	ATTTATAGTT TGTTCCTTTT	1020 1080
55	SEQ ID NO:222 Protein Accession	PC14 Protein sequ n #:	ence: NP_057654				
	1	11	21	31	41	51	
60	KYEYEVDKDF KEWQRMLQLI VAGNFHITVG	SLVKELDAFP SSKLRINIDI QSRLQEEHSL KAIPHPRGHA ITVVPTKLHT	TVAMKCQYVG QDVIFKSAFK HLAALVNHES	ADVLDLAETM STSTALPPRE YNFSHRIDHL	VASADGLVYE DDSSQSPNAC SFGELVPAII	PTVFDLSPQQ RIHGHLYVNK NPLDGTEKIA	60 120 180 240 300
65		FWQFFVRLCG					360
				SE	Q ID NO:223 PEZ:	B DNA SEQUENCE	
70	Nucleic Acid Acc Coding sequence		NM_001935.1 76-2301 (unde			and stop codons)	
75	1	11	21	31	41	51	
80	GAGGAGACGC GCGCTTGTCA ACAGCTGACA	CGCCGCCCGC CGACGATGAA CCATCATCAC GTCGCAAAAC CCTTAAGATG	GACACCGTGG CGTGCCCGTG TTACACTCTA	AAGATTCTTC GTTCTGCTGA ACTGATTACT	TGGGACTGCT ACAAAGGCAC TAAAAAATAC	GGGTGCTGCT AGATGATGCT TTATAGACTG	60 120 180 240 300
-	.um. manut		ween a worder!		20		

		TCAATGCTGA					360	
		GACATTCTAT					420	
		ACTACGTGAA					480	
5		AAAGGCAGCT					540	
,		CAGTGGGTCA ATTTACCAAG					600 660	
		CTGACTGGGT					720	
		ACGGCACTTT					780	
		CCTTCTACTC					840	
10		CAGGAGCTGT					900	
10		TCACCAATGC					960	
		ACTTGTGTGA					1020	
		TTCAGAACTA					1080	
		GCTTAGTGGC					1140	
15	AGATTTAGGC	CTTCAGAACC	TCATTTTACC	CTTGATGGTA	ATAGCTTCTA	CAAGATCATC	1200	
	AGCAATGAAG	AAGGTTACAG	ACACATTTGC	TATTTCCAAA	TAGATAAAAA	AGACTGCACA	1260	
	TTTATTACAA	AAGGCACCTG	GGAAGTCATC	GGGATAGAAG	CTCTAACCAG	TGATTATCTA	1320	
	TACTACATTA	GTAATGAATA	TAAAGGAATG	CCAGGAGGAA	GGAATCTTTA	TAAAATCCAA	1380	
00		ATACAAAAGT					1440	
20		TGTCATTCAG					1500	
		TCTATACTCT					1560	
		CTTTGGATAA					1620	
		TGAATGAAAC					1680	
25		AATATCCTCT					1740	
43		TCAGACTGAA					1800	
		ATGGCAGAGG					1860	
		GAACATTTGA					1920 1980	
		TCCTGGGATC				AGGGTACGTA	2040	
30		GGGAGTACTA					2100	
50		ACCTTGACCA					2160	
		AGTACCTCCT					2220	
		TCTCCAAAGC					2280	
		ACCATGGAAT					2340	
35		TAAAACAATG					2400	
	AAGCTTATTA	AAACTCATTT	TIGITITCAT	TATCTCAAAA	CTGCACTGTC	AAGATGATGA	2460	
		AATACACACT					2520	
		TCTTAAGTAG					2580	
40		CGGTCGGGTT					2640	
40		TTGTTTTTAT					2700	
		GACTGGTTCA CTAGGGCAGG					2760 2820	
		CCCAAGTCCA					2880	
		TTCACCACGA					2940	
45		AATATCGAAA					3000	
		AAGGGAAACT					3060	
		ACACAGCAAG					3120	
	TACTGATGTT	CCTAGTGAAA	GAGGCAGCTT	GAAACTGAGA	TGTGAACACA	TCAGCTTGCC	3180	
<b>5</b> 0	CTGTTAAAAG	ATGAAAATAT	TTGTATCACA	AATCTTAACT	TGAAGGAGTC	CTTGCATCAA	3240	
50		TTCATTTCTT					3300	
						ATTCTACATA	3360	
	CTATGGAATT	TCTCCCAGTC	ATTTAATAAA	TGTGCCTTCA	TTTTTTC			
55	SEU ID NO-334 I	PEZ3 Protein segu	ionco.					
<i>JJ</i>	Protein Accession		NP_001926.1					
	T TO COLL PAGE CONTROL		M _0010E01					
	1	11	21	31	41	51		
	1	1	1	1	1	i		
60	MKTPWKILLG	LLGAAALVTI	ITVPVVLLNK	GTDDATADSR	KTYTLTDYLK	NTYRLKLYSL	60	
	RWISDHEYLY	KQENNILVFN	AEYGNSSVFL	ENSTFDEFGH	SINDYSISPD	GQFILLEYNY	120	
	VKQWRHSYTA	SYDIYDLNKR	QLITEERIPN	NTQWVTWSPV	GHKLAYVWNN	DIYVKIEPNL	180	
		EDIIYNGITD					240	
65		TVRVPYPKAG					300	
65		ISLQWLRRIQ					360	
		FYKIISNEEG					420	
		LYKIQLIDYT					480	
		LRVLEDNSAL					540	
70		CSQKADTVFR					600 660	
70		RQFSKMGFVD MGLPTPEDNL					720	
		QAMWYTDEDH				ATT GOSAGIS	120	
	IAMVDIGUE.	g	GIMDOIIM.	11111111111111111111111111111111111111	40. 0			
				SE	2 ID NO:225 PBJ	DNA SEQUENCE		
75	Nucleic Acid Accession #: Coding sequence:		none found					
			1-261 (underlined sequences correspond to start and stop codons)					
	•	11	21	21	41	51		
80	1	ī	21 	31	ī	Ĭ		
	1	•	•		1	•		
					39	3		

5	AGAAGTGTGA AGAAACCCTC GTGGAAATGA	TTAAAGTGCG AGGAGCTCTG	GGAGCCAAAC TGCTAACCAG GATGGGCCTG AGTCTGGTCC A	TGTTCCCTGC CTCCTCTTGA	ATGAGGCAGA TGGGGGTCCT	AAGTGAATCC AGAAGCATGT	60 120 180 240
10	Protein Accession		none fo				
	•	11     ANDNAIRVON 'S LRDDKEQSF	21     RSVIKVRANQ PH OPTLDV	31     CSLHEAESES	41   RNPQELWMGL	51    LLLMGVLEAC	60
15		•		SE	O ID NO:227 PEM	2 DNA SEQUENCE	i
	Nucleic Acid Acc Coding sequence		none fo 1-462 (		ces correspond to	start and stop code	ens)
20							
25	CTCATACTTY ATTGATGTAY ATCATGTGG TTTATGGCT	G CTGTATGTTY T CTTCTCAAG A CCAGTTTTG A TTGAAGAAG	G TGGATCAGC. A TCTGGACAG. T GGAAGACAA A AATGAAGAA	A AATATAGTC: A CGGCCAGAG: I CTTTCCATG G CACGGAAGT:	A GCCCTCTAC: A GTATGCTGT: G GCTGGGGGAI A CTCATGTGG	51     AAAAACTGCT   TGAGCAAAAT   TCTAGTCATC   AGCTAGAAGAT   ATTCCCAGAA   TCCAAGGAAG	120 180 240 300
30	AGCAGAACA	C CTGAAAGCC		T GACACTGAG	A ATGAAGAGT	A TCACAGGTTT	
	SEQ ID NO:228 Protein Accession	PBM2 Protein seq n #:	<u>uence:</u> none fo	und			
35		٠					
	1	11 	21 	31 	41 !	51 {	
40	IMWISTVEDN		LILAVCCGSA FMAIEEEMKK				60 120
40	SRTPESQQFP	DTENEEYHRF	VKDQIVVDM	R RYF			
40					Q ID NO:229 PEZ:	2 DNA SEQUENCE	
40	SRTPESQQFP  Nucleic Acid Acc Coding sequence	ession#:	NM_014253	SE		2 DNA SEQUENCE and stop codons)	
	Nucleic Acid Acc	ession#:	NM_014253	SE			
	Nucleic Acid Acc Coding sequence 1     GACTGCTTGC	ession #: :: 11   ATTAAAGGAC	NM_014253 65-8242 (unde	SE  ifined sequences of the sequences of	correspond to stand 41   GAAACTGAGC	and stop codons) 51   TTGCTTAATC	60 120
45	Nucleic Acid Acc Coding sequence 1     GACTGCTTGC AGAGATGGAG GGATCTAGCT	ession #:  11  ATTAAAGGAC CAAACTGACT TACACCAGTT	NM_014253 65-8242 (under 21   TTCCTCATCC GCARACCCTA CTTCTGATGA	SE  idined sequences  31  TTTTTTCAT  CCAGCCTCTA  GAGTGAAGAT	correspond to stand  41    GARACTGAGC CCARAGTCA GGAAGAAAAC	and stop codons)  51    TTGCTTAATC AGCATGAAAT CAAGACAGTC	60 120 180
45	Nucleic Acid Acc Coding sequence 1     GACTGCTTGC AGAGATGGAG GGATCTAGCT ATACAACTCC CCAGAGTAGA	ession #:  11   ATTAAAGGAC CAAACTGACT TACACCAGT TACACCAGT AGGGAGACCC AAGAGGAAAG	NM_014253 65-8242 (under 21   TTCCTCATCC GCAAACCCTA CTTCTGATGA TGCACGAGTA ARGTAGAAAA	serifined sequences of the sequences of	41 GARACTGAGC CCARAGGTCA GGARGAAAC CTGAGGATGA GAGATGAAAC	and stop codons)  51    TIGCTTAATC AGCATGAAAT CAAGACAGTC ATTACAATAG TCTGTGAAAC	60 120 180 240 300
45	Nucleic Acid Acc Coding sequence 1 GACTGCTTGC AGAGATGGAG GGATTAGCT ATACAACTCC CCACAGTAGA CTCTCACACT CCACCTAGAG	ession #:  11   ATTANAGGAC CANACTGACT TACACCAGTT AGGGAGACAC CAGGGAGACAC CAGGGAGACAC ATGGGATCTG	NM_014253 65-8242 (under 21   TTCCTCATCC GCAAACCCTA CTTCTGATGA TGCACGAGTA AAGTAGAAAA GCTACCAAAC	strined sequences of the sequences of th	41   GAAACTGAGC CCAAAAGTCA GGAAGAAAAC CTGAGGATGA GAGATGGAAT AGCGTTTCTC GGTGCTGCCT	and stop codons)  51    TTGCTTAATC AGCATGAAAT CAAGACAGTC ATTACAATAG TCTGTGAAAC CGCATGGCTA CACCTGACCA	60 120 180 240 300 360 420
<b>45 50</b>	Nucleic Acid Acc Coding sequence 1   GACTGCTTGC AGAGATGGAG GGATCTAGCT ATACAACTCC CCAGAGTAGA CTCTCACACT CCAGCTAGAG TGCACTAAGA	ession #:  11   ATTAAAGGAC CAAACTGACT TACACCAGTT AGGGAGACCC AAGAGGAAAC CTGTGCTCTC ATGGGATCTG ATGGGATCAA	NM_014253 65-8242 (under 21   TTCCTCATCC GCARACCCTA CTTCTGATGA TGCACGAGTA TGCACGAGTA AAGTAGAAAA GCTACCAAAC	strined sequences of the sequences of th	41 GAAACTGAGC CCAAAAGTCA GGAAGAAAAC CTGAGGATGA AGCGTTCTC AGCTCCCCT AGCTCCCTGTT	and stop codons)  51   TTGCTTAATC AGCATGAAAT CAAGACAGTC ATTACAATAG TCTGTGAAAC GGCATGCCTA TGTCCAGCCG	60 120 180 240 300 360
<b>45 50</b>	Nucleic Acid Acc Coding sequence 1     GACTGCTTGC AGAGATOGAG GGATCTAGCT ATACAACTCC CCAGCTAGAG CTCTCACACT CCAGCTAGAG TGCACTAGAG GGCCAACTCT TGGTTTCAAA	ession#:  11  ATTANAGGAC CANACTGACT TACACCAGTT TACACCAGTT AGGGAGACC AAGAGGAAAC CTGTGCTCTG ATGGGATATC ATGTGGATATCT TTCTCTCTG	NM_014253 65-8242 (under 21   TTCCTCATCC GCAAACCCTA TGCACGAGTA AAGTAGAAAA GCTACCAAAC ATGTGGACAC TTGTGGACAC TTGTGTGACAC TTTGTTGTGACAC TTTGTTGTGACAC	serifined sequences of the sequence of the	41 GAAACTGAGC CCAAAAGTCA GGAAGAAAAC CTGAGGATGA AGCGTTTCTC GGTGCTGCCT AGTTCCTGTT AGGAAGTCTGC CAAGCTGGGT	and stop codons)  51   TTGCTTAATC AGCATGAAAT CAAGACAGTC ATTACAATAG TCTGTAAAC GCCATGCCTA CACCTGACCA TGTCCAGCCG ATGCCGAAAA CTACTCAAGA CTACTCAAGA	60 120 180 240 300 360 420 480 540 600
<b>45 50</b>	Nucleic Acid Acc Coding sequence 1   GACTGCTTGC AGAGATGGAG GGATCTAGCT ATACAACTCC CCAGAGTAGA CTCTCACACT CCAGCTAGAG GGCCAACTCT TGGTTTCAAA TGGCTTCAAGAC TGCATGAGC	ession #:  11  ATTAAAGGAC CAAACTGACT TACACCAGTT AGGGAGACCC AAGAGGAAAC CTGTGCTCTC ATGGGATCTG ATGTGATAA GCATTATCCT TTCTCTCCTG TGCACCACA TGCACCTGTG	NM_014253 65-8242 (under 21   TTCCTCATCC GCARACCCTA ACTACGAGTA AAGTAGAAAA CCTACCAAAC GCGGAATGAA TGACTGACAC TTTGTTGGACAC CCAGGTACCAC CCAGGAAGCC CCAGGAAGCC	strined sequences of the sequences of th	41 GAAACTGAGC CCAAAAGTCA GGAAGAAAAC CTGAGGATGA AGCGTTTCTC GGTGCTGCCT AGTTCCTGTT AGGAAGTCTG CAAGCTGGGT CTCCACCGC GCGGACTCTC	and stop codons)  51    TTGCTTAATC AGCATGAAAT CAAGACAGTC ATTACAATAG TCTGTGAAAC GGCATGACTA TGTCCAGCCG ATGGGGAAAA CTACTCAAGA CTACTCAAGA TCTCCAGCCG TTCAGAGGAG	60 120 180 240 300 420 480 540 600 720
45 50 55	Nucleic Acid Acc Coding sequence 1   GACTGCTTGC AGAGATGGAG GGATCTAGCT ATACAACTCC CCAGAGTAGA CTCTCACACT CCAGCTAGAG GGCCAACTCT TGGTTCAAGA TGTGCAGAGC TCCTCATGCC ATCAATGAC	ession #:  11  ATTAAAGGAC CAAACTGACT TACACCAGTT AGGAGACCC AAGAGGAAAG CTGTGCTCTG ATGGGATTAA GCATTATCCT TTCTCTCTG AGCCCACACA ACCCCACACA	NM_014253 65-8242 (under 21   TCCTCATCC GCAAACCCTA CTTCTGATGA TGCACGAGTA AAGTAGAAAC ATGTAGAAAC ATGTAGAAAC TGACTAACC GGGGAATGAA TGACTGACAA TGACTGACAA TGACTGACAA	serifined sequences of the sequences of	41  GARACTERGE CCARARGTCA GGARACTERGE GGARGARA AGCGTTTCTC GGTGCTGCTT AGGARGTCA AGCTGCTT AGGARGTCA CCARCGC CCCCCCCA GCTCCCCCAA	and stop codons)  51    TEGCTTAATC AGCATGAAAT CAAGACAGTC ATTACAATAG ATTACAATAG ACCTGACCA ATGCCGC ATGGGAAAA CTACTCAAGA CACCTCCGCC TTCAAGAG CACCTCCGCC TTCAAGAGAG CACCTCACCAC	60 120 240 300 360 420 480 540 600
45 50 55	Nucleic Acid Acc Coding sequence  1	ession #:  11    ATTANAGAC CANACTGACT TACACCAGTT AGGGAGACAC AGGGAGACC ATGGGATAT CTGTGCTCTG ATGTGGATAA GCATTATCCT TTCTTCTCT TCTCTCTCT AGCCCACACA TGCACCTGTG ACCCGCAGCC CATCTGCATA	NM_014253 65-8242 (under 21	stimed sequences of the	41   GAAACTGAGC CCAAAAGTCA GGAAGAAAAC GGAAGAAAAC GGAAGATGAAA AGGATTCTC GGTGCTGCT AGTACTGTT AGGAAGCTGT CTACCACCGC GCGGACTCTC GCTCCCCACCGC GCTCCCCCACTT TTCAGTGCAG	and stop codons)  51    TTGCTTAATC AGCATGAAAT CAAGACAGTC ATTACAATAG GCCATGCCTA CACCTGACCA TGTCCAGCCG ATGGGGAAAA CACCTCCACC TTCAAGAG CACCTCAGCA TGTCAGCAG CAGCACGCA TGGAGCAG CCAGCACGAA	60 120 180 240 300 420 480 540 660 720 780 840 900
45 50 55	Nucleic Acid Acc Coding sequence 1   GACTGCTTGC AGAGATGGAG GGATCTAGCT ATACAACTC CCAGAGTAGA CTCTCACACT TGGTTCAAACT TGGTTCAAA TGTGCAGAGC TCCTCATGCC ATCAATGACT GGATTCACT GGATTCACT GGATTCACT CTACCCTCTG CTACCCTCTG CTTCCCCGA	ession #:  11  ATTAAAGGAC CAAACTGACT TACACCAGTT AGGAGGACAC AMGAGGAAAG CTGTGCTCTG ATCTGCATTATCCT TTCTCTCTG AGCCCACACA ACCACCTGTG ACCCGCAGCC CATCTGCATA TCAAACATA CCTGCCTTTA	NM_014253 65-8242 (under 21	serifined sequences of the sequences of	41   GARACTERGE CCARARGTCA GGARGATGA GGAGATGA AGCGTTTCTC GGTGCTTCTCTT AGGARGTCA CTCCACCGC CCGGGACTCTC CCCCCCAA AACATACCAT TTCAGTGCAG AGCCTCTTCCCTTT TGCTGCACCT TTCCACCTCTT TCCCACCTCTT TCCCACCTCTT TCCCACCTCTT TCCCACCTCTT	and slop codons)  51    TTGCTTAATC AGCATGAAAT CAAGACAGTC ATTACAATAG CGCATGGCTA ACTGCCGC ATGGGGAAAA CTACTCAAGA CTACTCAAGA CTACTCAAGA CTACTCAGCG ATGGGGAAAA CTACTCAGCG CCAGCACGCA TGGAGAGCAGCA TGGAGACCAG CCAGTCAGAA CTGCAAGCAC GGAAGTCCAC	60 120 180 240 360 420 550 600 660 720 780 840 900 960 1020
45 50 55 60	Nucleic Acid Acc Coding sequence  1	ession #:  11   ATTANAGGAC CANACTGACT TACACCAGTT AGGGAGACAC ATGGGATCTG ATGGGATCTG ATGGGATCAC ATGCACTGTG ACCCACACA TCCACCTGTG ACCCGCAGCC CATCTGCATA TCAAACATG ACATCCATT TCAAACATG ACATCCATTA ACATCCAATA ACATCCAATA CCTGCCTTTA GCCACTGCAA	NM_014253 65-8242 (under 21   TTCCTCATCC GCARACCCTA CTTCTGATGA AGTAGAAAA ATGTGGACAC ATGTGGACAC ATGTGGACAC ATGTGGACAC CTTTTGTGA ACCAGTTCAC ACCAGTTCAC ACCAGTTCAC CCAGGAAGCC ACAGCTGGGT GATCTGGTT CCGTTGACTA TCACAGTGACA TCACAGTGACA TCACAGTGACA	serimed sequences of the sequences of th	41   GAAACTGAGC CCAAAAGTCA GGAAGAAAAC CTGAGGATGA AGCGTTTCTC GGTGCTGCT AGTTCCTGTT AGGAAGTCA CCGAAGTCAC CCGACGC CCCCCA AACATACCAT TTCAGTCAG TTCAGTCAGC TTCAGTCACT TTCAGTCAGC TTCAGTCAC TTCAGTCAC TTCAGTCAC TTCAGTCAC TTCAGTCAC	and stop codons)  51    TTGCTTAATC AGCATGAAAT CAAGACAGTC ATTACAATAG TCTGTAAAC GGCATGCCTA CACCTGACCA TGTCCAGCCG TTCAGAGGAAA CACCTCAGAC CCAGCACCAG CCAGCACCAG CCAGCACCAG CCAGCACCAG CCAGTCAGAA CTCGAAGCAC ATGTGATTCC ATGTGATTCC	60 120 180 240 300 360 420 480 540 660 720 840 900 960 1020 1080
45 50 55 60	Nucleic Acid Acc Coding sequence  1	ession #:  11    ATTANAGGAC CANACTGACT TACACCAGTT AGGGAGACCC AAGAGGANAG CTGTGCTCTG ATGGGATCTG ATGTGCTCTG ATGCCACACA TCCACCACACA AGCCCACACA ACCCGCAGCC CATCTGCATA ACCACTGTG ACCTCTTA ACCACTGTA ACATCCAATA CCTGCCTTTA ACCACTGCATA ACACCACTGCA ANAGGGAACA ANAGGGAACA	NM_014253 65-8242 (under 21   TTCCTCATCC GCAAACCCTA ACTTGATGA TCCACGAGTA AAGTAGAAAA CCTTACCAAAC GGGGAATGAA TGACTGACAC CCAGGAAGCA CCAGGTCAC ACAGCTGGC ACAGCTGGC CCAGCTGGTC CCTTTAACAA TCACAGTGA TCACAGTGAC CCTTGACAA TCACAGTGAC CGGGGACCGA	stimed sequences of the	41  GARACTGAGC CCAAAAGTCA GGAAGAAAAC CTGAGGATGA GAGATGGAAT AGCGTTTCTC GGTGCTGCT AGGAGCTGGCT AGGACTGGT AGCACTGC CCCCCAA AACATACCAT TTCAGTGCAG AGGCTCTTC TGCTGCACCT TGCTGCACCT TTACATGCAG AGCACTTTC TGCTGCACT TTACTAGCAC TAACTACAT TCAGTGCAG AGCACTCTTC TGAGGAGC AACATACTACT	and slop codons)  51    TIGCTTAATC AGCATGAAAT CAAGACAGTC ATTACAATAG TCTGTGAAAC GGCATGACA TGTCAACCA ATGTCAACCA ATGTCAACCA ATGTCAACCA ATGTCAACCA CACCTCCACCC TTCAGAGGAG CCAGCACGCA TGGAGACCAG CCAGCACGCA TGGAGACCAC GGAAGTCAGAA CTCGAACCAC GGAAGTCACA ATGTGATTGC TGTATGCAAA CTCCAAATTGG	60 120 180 240 300 360 420 540 660 660 720 780 840 900 960 1020 1020 1140 1200
45 50 55 60	Nucleic Acid Acc Coding sequence  1	ession #:  11    ATTANAGGAC CANACTGACT TACACCAGTT AGGGAGANAG CTGTGCTCTG ATGTGATAA TTCACTG ACCCACACA TCCACCTGTG ACCCGCAGCC CATCTGCATA ACTGCATTA TCCACTGTGATAAACATG ACATCCATT TCCACTGCATA TCCACTGCATA TCCACTGCATA TCCACTGCATA TCCACTGCATA TTCAACATG ACATCCATA TTCAACATG ACATCCATA TTCGCCTTTA TCCGCCTGA AAAGGGAACA TTCGGCCTGA TTCGGCCTGA TTCGGCCTGA TTCGGCCTGA TTCGGCCTGA TCTGATAAAT GACATTGGTG	NM_014253 65-8242 (under 21   TTCCTCATCC GCAAACCCTA TGCACGGGATA AAGTAGAAAA ATTGGACAC GGGGAATGAA ACCAGTTCAC CCAGGAGAC ACCAGTTCAC CCAGGAGAC ACCAGTTCAC CCAGGAGAC CCTGTTACTC CCTTTAACAA TCACAGTGAC TCTGGCAGT CCTTGGCAGT CCTTGGCAGT CCGGGACCAG CCAGGGAACAC CACAGGAAAAA CACAGGTCAT	serimed sequences of the sequences of th	41   GAAACTGAGC GGAAGAAAAC CTGAGGATGA AGCGTTTCTC GGTGCTGCT AGTTCCTGTT AGTAGCACGC CCGAAGAGTCA AGCATCTAC CAAGCTGGT TTCCACCGC AACATACCAT TTCAGTGCAG AGCACTCT TTCAGTGCAG CAAGCACT TTCAGTGCAC CAAGCACT TTACTAGCCT GAAGGAGAGC ACTACTACT AAGGAAGAGC CCACCTGGTT	and slop codons)  51    TTGCTTAATC AGCATGAAAT CAAGACAGTC ATTACAATAG CTCTGCAAC GGCATGCCTA CACCTGACCA TGTCCAGCGG ATGCTCAAGA CACCTCAGAGAAA CTACTCAGAG CCAGCACCA TGGAGACCAG CCAGTCAGAA CTCGAGCAC ATGTGATTGC TGTATGCAAA CTCCAATTGG CGATAGACAC TGTATGCAAC TGTATGCACC TATTCTGCCG TATTCTGCCG TATTCTGCCG TATTCTGCCG TTTCTGCCG TTTCTGCCG TTTCTGCCG TTTCTGCCG TTTCTGCCG TTTCTGCCG TTTTCTGCCG TTTTCTGCCG TTTTCTGCCG TTTTTCTGCCG TTTTTTTT	60 120 180 240 300 360 420 480 540 600 660 720 960 900 960 1020 1080 1140 1200 1260 1320
45 50 55 60 65	Nucleic Acid Acc Coding sequence  1	ession #:  11    ATTANAGAC CANACTGACT TACACCAGTT AGGGAGACC AMAGGANAG CTGTGCTCTG ATGTGGATAA GCATTATCCT TTCTCCTG ACCCGCAGCC CATCACAGCA ACCCGCAGCC CATCACATA ACCACTGCTTA ACCTGCCTTTA ACCACTGCATA TCCACTGCAACA TCTGCCTTGAACAT GCACTGGAACA TCTGATAACA TCTGATAACA TCTGATAACA TCTGATACAC ACATTGGTG ACATTCACC ACTATCCACC ACTATCACC	NM_014253 65-8242 (under 21	stimed sequences of the	41   GAAACTGAGC CCAAAAGTCA GGAAGAAAAC GGAAGAAAAAC CTGAGGATGAA AGGATTCATC GGTGCTGCT AGTACTGTT AGGAAGTCTT AGGAAGTCTT AGGAAGTCTT CAACCTGGT TTCCCCACCGC GCGGACTCTC GCTCCCCAAA AGGACTGTT TTCAGTGCAG AGGCCTCTTC TGCTGCAACT TTAACTAGCCT GAAGGAGGAC ACTACTTACTT AAGGAACGT CCACCTGGTT AATATTTCTT	and slop codons)  51    TTGCTTAATC AGCATGAAAT CAAGACAGTC ATTACAATAG TCTGTGAAAC GGCATGCTA TGCTGCAAC TGTCAGCCA TGTCAGCAG ATGGGAAAA CTACTCAAGA CACTCAGCAG CCAGCACGCA CCAGCACGCA CCAGCACGCA CTGAAGCAC CCAGTCAGAA CTCGAAGCAC GGAAGTCCAC TGTATGCAAA CTCCAATTGG CGATGAAA CTCCAATTGG CGATGAACAC TATTCTGCGC TATCTGGCG TAGCCAAGGA TAGCCAAGGA	60 120 180 240 300 420 480 540 660 720 780 900 900 91080 1140 1200 1220 1320 1380
45 50 55 60 65	Nucleic Acid Acc Coding sequence  1	ession #:  11    ATTANAGGAC CANACTGACT TACACCAGTT AGGGAGACCC AAGAGGARAC CTGTGGATAT ACGTTATCT ATTCTCTCTG AGCCACACA TCCACTGTG ACCCGCAGCC CATCTGCATA TCAACACTGTG ACACTCCATA TCAACACTGAC TCCACTCAA TTCAACACTGAC TCCACTCAA TCGGCCTGA AAAGGGAAC ATTGGTCAA TCTGATAAAT GACATTGGTG ACATTGGTG ACATTGGTG ACATTGGTG ACATTGGTG ACATTGATAAAT GACATTGTG ACTATCCACC GGAATTTATG ATGGATGGCA	NM_014253 65-8242 (under 21   TTCCTCATCC GCARACCCTA CTTCTGATGA AGTAGAAAA ATGTGGACAC ATGTGGACAC ATGTGGACAC ATGTGGACAC CCAGGAGTA ACCAGTTCAC ACCAGGAGAC ACCAGGCC ACAGCTGGGT GATCTGGTTC CCTTTAACAA TCACAGTGAC CCTGGCAC CCTTGCCAG	serimed sequences of the sequence of the	41   GARACTGAGC CCAAAAGTCA GGAAGATGA GGAGATGA AGGATGA AGGATGT AGTTCCTGT AGTTCCTGT AGTTCCTGT AGTTCCTGT AGTACTGCT AGTACTGCT AGTACTGCT CTACACGC GCGGACTCT CTCCACCGC AACATACCAT TTCAGTGCAG AGCCTCTTC TACTAGCACT TACTAGCT AAGGAGAGC CACTGCTACT AAGGAGAGC CCACTGCTT AATATTTCTT ACACATACTC TCCAAGGCCT TCCAAGGCCT TCCAAGGCCT CCCAAGGCCTCTC CAAGGACGGC CCACTGGTT AATATTTCTT TCCAAGGCCT TCCAAGCCT TCCAAGCT TCCAAGCCT TCCAAGCT TCCAAGCCT TCCAAGCCT TCCAAGCT TCCAAGCT TCCAAGCCT TCCAAGCT TCCA	and slop codons)  51    TTGCTTAATC AGCATGAAAT CAAGACAGTC ATTACAATAG CGCATGGCTA CACCTGACCA TGTCCAGCCG TCACCACA TCACCAGCA CCACCACCA TCACCAGCAC TCACCACCA TCACCACCA TCACCACCA TCACCACCA TCACCACCA TCACCACCA TCACCACCA TCACCACCA TCGACCAC CCACTCAGAA CTCAATTGC CGATATGCAC TCTATCCAATTGC TCGATCAACAC TATTCTGCCG TAGCCAAGGA AGTTTTGATTTC TTGATTTATCATAC	60 120 180 240 360 420 480 5540 660 720 780 840 900 1020 1140 1260 1320 1320 1340 1500
45 50 55 60 65	Nucleic Acid Acc Coding sequence  1	ession #:  11    ATTANAGGAC CANACTGACT TACACCAGTT AGGGAGACAC AGGGAGACAC ATGGGATATACCT ATGTGGATAA TCCACCACACA TACACCACACA TACACCACACA TACACCACACA TACACCACACA TACACCACACA TACACCACACA TACACCACACAC TACACCACACA TACACCACACA TACACCACACA TACACCACACA TACACACAC	NM_014253 65-8242 (under 21	serimed sequences of the sequence of the	41   GAAACTGAGC CCAAAAGTCA GGAAGAAAAC CTGAGGATGA AGCGTTTCTC GGTGCTGCT AGTTCCTGTT AGGAAGTCA AGGAAGTCA AGGAAGTCA AGGAAGTCA TTCAGTCAC AGCCCCAA AACATACCAT TTACTAGCCT GAAGGAGCC CTACTACTAC AAGGAGAGC CACTGGTT AATATTCTT ACACATACTT ACACATACTT CCAAGGGCT CCAAGGGCT CCAAGGGCT CCAAGGGCT CCAAGGGCT CCAAGGGCT CCAAGGGCT CAAGAGCCT CCAAGGGCT CAAGAGCCT CCAAGGGCT CAAGAGCCT CAAGAGCCT CAAGAGCCT CAAGAGCT CAAAGAGCT CAAAGAGCT CAAACAGGCT CAAAACAGGCT CAAACAGGCT CAAACAGACAC CAAAACACAGCT CAAACACACAC CAAACACACAC CAAACACACACC CAAACACACC CAAACACACACC CAAACACACACC CAAACACACC CAAACACACACC CAAACACACACC CAAACACACACC CAAACACACACC CAAACACACACC CAAACACACACC CAAACACACACC CAAACACACACACC CAAACACACACC CAAACACACACC CAAACACACACC CAAACACACACC CAAACACACACC CAAACACACACACC CAAACACACACACC CAAACACACACACC CAAACACACACACC CAAACACACACC CAAACACACACACACC CAAACACACACACACC CAAACACACACACC CAAACACACACACACACACACACACACACACACACACACA	and slop codons)  51    TTGCTTAATC AGCATGAAAT CAAGACAGTC ATTACATTAG TCTGTGAAAC GCCTGACCA ATGCCAGCCA ATGCGGAAAA CTACTCAAGA CACCTCAGCC ATGGGAAAA CTACTCAAGA CCACTCAGCA TGGAGACCAG GGAGACCAG GGAGTCAGAA CTGGAAGCAC GGAAGTCAGAA CTCCAATTGG CGATAGGAA CTCCAATTGG CGATAGGAA CTCCAATTGG CGATAGGAA AGTTTCATTT	60 120 180 240 300 360 420 480 540 660 720 840 900 960 1020 1140 1200 1320 1380 1440 1500 1560
45 50 55 60 65 70	Nucleic Acid Acc Coding sequence  1	ession #:  11  ATTAAAGGAC CAAACTGACT TACACCAGTT AGGGAGACCC AAGAGGAAAG CTGTGCTCTG ATGGGATCTG ATGGGATCA AGCACACA TCCACCACACA AGCCCACACA ACCCGCAGCC CATCTGCATA ACCACTGATA ACATCGATA ACATCGATA ACATCGATA ACAGCGAACA TCGACTGATA ACATCGATA ATGGCCTAAAAAT CCTGCCTTTA GCACTCCAA AAAGGGAACA TCTGATAAAT GACATTGGTG ACATTCACC GGAACTTGTG ACTATCACC GGAACTTGGTA ACTGCTCGGAACC CCTCGGAACC CCTCGGAACC CGGACCTTGGTA ACTACAGCAA	NM_014253 65-8242 (under 21 TTCCTCATCC GCAAACCCTA TCCTCATCC GCAAACCCTA AAGTAGAAAA TCACAGAGTA AAGTAGAAAA TCACTGACAC CCAGGAATGAA TCACTGACAC CCAGGAAGCC ACAGCTCGC CCTTTAACAA TCACAGTGAC CCTGGCACT CCTTTAACAA TCACAGTGAC CAGGAAAAA CACAGTCGAT CACAGTGAC CAGGAAAAA CACAGGTCAT TCACAATATA GCAGAAGAAA AACAGCTCGT TGAAATAAT TCACAATATA TCACGAGTCAT TTGAAATAAT	serifined sequences of the sequences of	41  GARACTERGE CCARARGTCA GGARACTERGE GGARACTERGE GGARGARAC GGARGARAC AGCOTTTCTC GGTGCTGCTT AGGARGCTGGCT AGCACGGC CCCCCAR ARCATACCAC TCCACCGC GCTCCCCCCAR ARCATACCAC TTCAGTGCAC TTCAGTGCAC TTCAGTGCAC TTCAGTGCAC AGCGCTCTT CAAGGACAC TTCATACT TACTAGCCT TACTACT CAAGGACAC ACTACTTACT AAGGACAC ACTACTTACT AATATTCTT ACACATACTC CCAAGGGCT CCCAAGGCCT CCCAAGGCCT CCCAAGGCCT CCCAAGGCCT TCCAAGGGCT CCCAAGGCCT CCCAAGACAC TCCAACCCAATT	and slop codons)  51    TTGCTTAATC AGCATGAAAT CAAGACAGTC ATTACATTAG CTGTGAAAC GCCTGACCA ATGTCCAGCCG ATGGGGAAAA CTACTCAAGA CACCTCGCC TTCAGAGGAG CCAGCACGA TGGAGACCAG GGAAGTCAGA ATGTGATGAC GGAAGTCAGA ATGTGATTGC TGTATGCAAA CTCCAATTGG CGATAGAAA TTCTGAGG ATGTGATTTC TGATGATAC TAGTCAAGA AGTTTCATTT CTGATGATAC TCAATGAGAA TGGAGCAAGGA AGTTTGATTT CTGATGATAC TCAATGAAAA TCAATAGAAA TCAATAGAAA TCAATAGAAAA TCAATAGAAAA TCAATAGAAAA TCAATAGAAAA	60 120 180 240 300 360 420 6600 6600 6720 780 840 900 1020 1140 1200 1260 1320 1380 1440 1500 1500 1620 1680
45 50 55 60 65 70	Nucleic Acid Acc Coding sequence  1   GACTGCTTGC AGAGATGGAG GGATTAGCT ATACAACTCC CCAGAGTAGAG TCCACACTAGAG TGCACTAGAG TGCACTAGAG TGCACTAGAG TGCACTAGAG TGCACTAGAG TGCACTAGAG TGCACTAGAG TGCATTCACAT TGGTTTCAAA TGTTCAGAG CTTTTCCCGA AGCATTGAG AGCATTGAG AGCACTCT TGGAGAAAGTT TTGCAGAGT TCTCTCTGCTG TGTAAAACTA ACAGCACTCC TATGGATCAA ATTCGTGTTA ACAGCACTCC TATGGATCAA TTGGAGAGTT TGAGAGATTC	ession #:  11  ATTANAGEAC CANACTGACT TACACCAGTT AGGGAGACAC AGGAGACAC ATGGGATATA GCATTATCCT TTCTTCTTC TCTCTCTGAC ACCACACA TCCACCACAC TCCACCACACA TCCACCTTA ACCTCCATTA ACCTCCATTA ACCTCCATTA TCCACTGCATA TCCACTGCATA TCCACTGCATA TCGACTGCATA TCGACTGCATA TCGACTGCATA TCGACTGCATA TCGGCTTTA ACCTCCCAC CGAATTTATG ACTACCAC GGAATTTATG ACTACCAC GGACTTGGAC GGACCTTGGAC GGACCTTGGT ACTACAGCAA ACTCTCGCC TCCCCTGTC	NM_014253 65-8242 (under 21	seinined sequences of the sequence of th	41   GAAACTGAGC CCAAAAGTCA GGAAGAAAAC GGAAGATGA AGCGTTTCTC GGTGCTGCT AGTTCCTGTT AGTAGCAGCTGAGAAGTCA AGCAAGTCAC CCAACGC CCGCGCTCCCA AACATACCAT TTCAGTGCAG AACATACCAT TTACTAGCCT GAAGGAGCC CCACCTGGTT AATATTCTT ACACATACTA CCAACAGCCT CCAACGAGCT CCAACGAGCT CCAACGAGCT TCACTACTTACT CCAACGAGCT TCACAACAAT TCAAGCAGTT TCACAACAAAT TCAAACAAAT TCAACAAACA TCAACCAAC	and slop codons)  51    TIGCTTAATC AGCATGAAAT CAAGACAGTC ATTACAATAG TCTGTGAAAC GCATGGAAA CACCTGACCA ATGCGCAAAA CTACTCAAGA CACCTCCGCC TTCAGAGCAG CCAGCACGCA TGGAGACCAG CCAGTCAGAA CCTCGACCA TGGAGACCAG CCAGTCAGAA CTCGAAGCAC GGAAGTCCAC TGTATGCAAA CTCCAATTGG CGATAGAACAC TATTCTGGCG TAGCCAAGGA AGTTTGATTT CTGATGATAT CTGATGATAC TCATAGAGATA TGGAGCAAGT	60 120 180 240 300 360 420 480 540 660 720 840 900 960 1020 1140 1250 1320 1380 1440 1500 1560 1620 1620 1680

	AGGAGAAATA	TGCGAGGAAG	AGGACTGCCT	AGACCCAATG	TGTTCCAACC	ATGGCATCTG	1980
,	TGTAAAAGGA	GAATGTCACT	GTTCTACTGG	CTGGGGAGGA	GTTAACTCTC	AAACACCACT	2040
				CGGAACTTTT			2100
	CACCOTOTA A	CARGAGEAGI	GC TCAGGACA	COGARCIIII	CIICIGGACG	CIGGAGIAIG	
5						CCATGGAGTG	2160
3	TGGTAGCCAT	GGAGTCTGCT	CAAGAGGAAT	TTGCCAGTGT	GAAGAAGGCT	GGGTAGGACC	2220
	AACATGTGAG	GAACGCTCCT	GTCATTCTCA	TTGTACTGAG	CATGGCCAAT	GCAAAGATGG	2280
				CGACCACTGC			2340
				CTTTGGAAAT			2400
10	Aaatggttgg	CACTGTGTGT	GTCAGGTGGG	TTGGAGTGGG	ACAGGCTGCA	ATGTTGTCAT	2460
10	GGAAATGCTT	TGTGGAGATA	ACTTGGACAA	TGATGGAGAT	GGTTTAACCG	ACTICTICTICA	2520
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				TCTCTTCTCT			2640
	TTATGATCGA	ATCAAATTCC	TCATTGGCAA	GGACAGTACT	CATGTCATTC	CTCCTGAGGT	2700
	GTCATTTGAC	AGCAGGCGTG	CCTGTGTGAT	TCGAGGCCAA	GTGGTGGCCA	TAGATGGAAC	2760
15				GCACCACAGT			2820
10							
				CATCGGTGGC			2880
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	TGTGGTAGAG	AAAGTCACCA	TGCAGAGAGT	TGTATCAGAC	CCGCCATCCT	GCGATATCTC	3000
				TCCTTCACCG			3060
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20				GCTGCAGGTT			3120
	TCCCTCCAGC	TTTGTGAGGC	TGAGTTACCT	GAGCAGCCGC	ACCCCTGGGT	ATAAAACCCT	3180
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				GTGGTTTCCC			3300
				TGGACAGAAG			
25							3360
23	TTTGGTATCT	GTGGGATATG	AATATGAAAC	GTGCCCTGAC	TTTATTCTCT	GGGAGCAAAG	3420
	GACAGTCGTT	TTACAAGGTT	TTGAGATGGA	TGCTTCTAAC	CTAGGAGACT	GCTCTTTGAA	3480
				AATCATACAT			3540
				AACCATAATG			3600
20				CCACAACAAC			3660
30	CTTAGCTTCT	GGCCCTGATG	GCAGTGTGTA	TGTTGGCGAC	TTCAATTTTG	TAAGGAGAAT	3720
	ATTTCCCTCG	GGAAACTCCG	TTAGTATTTT	GGAATTAAGC	ACAAGTCCTG	CTCACAAATA	3780
				ACTCTATCTA			3840
				GAAAGATCTG			3900
م ج	GGCAGGAACT	GGTGATCAGT	GCCTTCCCTT	TGACCAGAGT	CATTGTGGAG	ATGGTGGGAG	3960
35	AGCATCGGAA	GCTTCACTGA	ATAGCCCTCG	AGGCATCACA	GTTGATAGGC	ATGGATTTAT	4020
				AATTGATGAG			4080
				ACCACTGAGC			4140
	CACTCAGGTG	CGATTAGAGT	GGCCAACAGA	CCTTGCAGTA	AATCCTATGG	ACAATTCATT	4200
	GTATGTCTTG	GATAACAACA	TTGTGCTGCA	AATTTCTGAG	AACAGGCGTG	TTCGGATCAT	4260
40				AGGCATCGAT			4320
				GGCCATCAGT			
							4380
				AAACCGCATT			4440
	GGAGATCTAC	ATCATCGCTG	GTGCCCCCAC	TGACTGTGAC	TGCAAAATTG	ATCCAAACTG	4500
				CAAAGATGCA			4560
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73				TGTGGCAGAC			4620
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	CGCTGATCAG	GAACTGTACC	AGTTCACTGT	AAATGGAACC	CACCTACACA	CCCTGAACTT	4740
	GATAACAAGG	GACTATGTTT	ATTACTTCAC	CTACAATTCT	GAAGGTGACT	TOCOCCO	4800
				TCGCCGTGAT			4860
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<b>50</b>				GCTGACTATA			4920
	AAGAGTGTCA	GCCCAAGGCT	ATAATCCGGC	CTTAATGACC	TATCCAGGAA	ACACAGGGCT	4980
	TCTGGCTACC	AAAAGTAACG	AAAATGGATG	GACAACCGTT	TATGAGTATG	ACCCCGAGGG	5040
				AGAGGTCAGC			5100
55				CAACCGTGAA			5160
<i>აა</i>				AAAACAAGAA			5220
	GGTGAATCCA	GATGGTTCCC	TGCGTGTCAC	TTTTGCCAGC	GGGATGGAGA	TCGGCCTCAG	5280
						GCAACATCTC.	5340
				CGAGTGGCGG			5400
<b>4</b> Λ				GAGGGCCCAC			5460
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						GAGGAACGTĜ	
				GAAAATTATT			5700
~~	GAAAATTTGG	AGCTATACCT	ACTTAGAAAA	ATCTGTGATG	CTTCTCCTAC	ACAGCCAGCG	5760
65	GCGTTACATC	TTTGAGTATG	ACCAATCAGA	TTGCCTGCTG	TCAGTTACCA	TGCCTAGCAT	5820
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				CTATAGTCGA			5940
				ATACAAGTAC			6000
	TGAGGTTCTC	TATGATACCA	CTCAGGTCAC	ATTAACATAT	GAAGAGTCTT	CTGGAGTGAT	6060
70				CATCTGCACA			6120
. •							
				CAGTGAAGAA			6180
				AAGCATGCAA			6240
	TTTGCCTATA	GATCTTTACC	GATATGTTGA	TGTCTCTGGC	AGAACAGAGC	AGTTTGGAAA	6300
				GGTCATAACT			6360
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, 5	CAAAATCTTC						6420
	TGCCTACTGG	ATGACCATTC	AATATGATAA	TGTGGGCCGA	CATGGTAATA	TGTGCATAAG	6480
						ATGGGCAACT	6540
	TCAGACTGTT						
							6600
oΛ						ATGACCTCCG	6660
80	AGACCGCATC	ACCAGATTAG	GAGAAATTCA	GTATAAAATG	GATGAAGATG	GCTTTCTGAG	6720

	GCAGAGGGGA	AATGATATTT	TTGAATATAA	TTCTAATGGC	CTGCTGCAGA	AAGCCTACAA	6780
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			ACCTTCAGTT				
_			ACACAAGCTC				
5	AGGTCACCTT	ATTGCCATGG	AGTTAAGCAG	TGGTGAAGAA	TATTATGTAG	CCTGTGATAA	7020
	TACAGGTACC	CCACTAGCTG	TGTTCAGCAG	CCGAGGTCAG	GTCATAAAGG	AGATACTATA	7080
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			TCCTTACTAA CAACGGCCTA				
10			ACTCCTTTGA				
10			ACATCAGAAG				
			CCAAACCTGA				
			CTCAAGAGTG				
			TCAGGAATTT				
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•	TTCTGTTTTT	GGGAAAGGTA	TAAAATTTGC	CATCAAGGAT	GGCATAGTAA	CAGCTGATAT	
	TATAGGAGTA	GCCAATGAAG	ATAGCAGGCG	GCTTGCTGCC	ATTCTCAATA	ATGCCCATTA	7740
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			ATGGAGCCCT				
			TGTTGGAGAT				
			TGCAAGAGGG				
25			GCACTGGGCG AACTTTCTGA				8160 8220
			AACAAAAATA				
			TTTATTGTAT				
			TCCAACTGCC				8400
			AAGAAATGAC				8460
30			TCAAACAGCT				
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			CTTAACTGTT				8640
			TGCCTTTCGA				8700
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			ACGACCTGTT				8940 9000
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			CGGTAATTTA				9180
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45	AAGGGAAGAC	CAGACCAAAC	ATCACAGCAG	TTGCTGCCAC	ATTGTTTCAG	CCCACTTAGA	9360
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			AGTTAGTTAT				9540
			CTAGGAAAAT				9600
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			TAGAAAATTC				9840
			TCTGAAGGCA				9900
			CAGTGTGGGG				9960
55			AAGCTCATGT				10020
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			CTCTCTTTAT				
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			GACCTTAAGT				
			CTGTATATAG CAATATTAGA				
			TGATGTATCA				
65			CTTAATAGTG				
			GTGCCTTTAG				
			TTACAGCTGT				
	TCTTTTTACA	ACTCCTAAAG	CTTGAGGGAG	GAAAGAAAAA	АААААСАААА	CTACTAATCA	10800
70			ATTTTGGCAT				
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			GAGCATAGTA				
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, ,	GCTTTTCTGT	ACCOUNT ACCOUNT	TTTCACTCTT	GCACTACAGT	CTAGAGATCC	AAATGAACTG	11220
	AMMAGITUAA TOTOLOGIA	CCD4CDC4CC	ATTTAAATAT TGAAGGAAAG	GETTACTET	AGTIGICATT	ACTIALICATIA	11240
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	AAAATAAGTY	TGTCCTTTAC	TGTCAATTTA	TCGAGAAGAT	CTATAATATA	TAGACTACAT	11460
80			AGCCAAATGT				
*		3-					

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        TTGGAGCATA TTATATATAG CTTGTGGAAA GACATAAGGC TACAGATGGA ATGGAACATT 11880
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        TATTTCATTT TTAAAAAATA AAATTGCCAA TGAAAAAAA
25
        SEQ ID NO:230 PEZ2 Protein sequence:
        Protein Accession #:
                                 NP_055068
30
                    11
                               21
                                           31
                                                                   51
        MEOTOCKPYO PLPKVKHEMD LAYTSSSDES EDGRKPROSY NSRETLHEYN OELRMNYNSO
                                                                                  60
        SRKRKEVEKS TOEMEFCETS HTLCSGYQTD MHSVSRHGYQ LEMGSDVDTE TEGAASPDHA
                                                                                 120
        LRMWIRGMKS EHSSCLSSRA NSALSLTDTD HERKSDGENG FKFSPVCCDM EAQAGSTQDV
35
        QSSPHNQFTF RPLPPPPPPP HACTCARKPP PAADSLQRRS MTTRSQPSPA APAPPTSTQD
        SVHLHNSWVL NSNIPLETRH SLFKHGSGSS AIPSAASQNY PLTSNTVYSP PPRPLPRSTF
                                                                                 300
        SRPAFTFNKP YRCCNWKCTA LSATAITVTL ALLLAYVIAV HLFGLTWQLQ FVEGELYANG
                                                                                 360
        VSKGNRGTES MDTTYSPIGG KVSDKSEKKV FQKGRAIDTG EVDIGAQVMQ TIPPGLFWRF
                                                                                 420
        QITIHHPIYL KFNISLAKDS LLGIYGRRNI PPTHTQFDFV KLMDGKQLVK QDSKGSDDTQ
HSPRNLILTS LQETGFIEYM DQGPWYLAFY NDGKKMEQVP VLTTAIEIMD DCSTNCNGNG
                                                                                 480
40
                                                                                 540
        ECISCHCHCF PGFLGPDCAR DSCPVLCGGN GEYEKGHCVC RHGWKGPECD VPEEQCIDPT
                                                                                 600
        CFGHGTCIMG VCICVPGYKG EICEEEDCLD PMCSNHGICV KGECHCSTGW GGVNCETPLP
        VCQEQCSGHG TFLLDAGVCS CDPKWTGSDC STELCTMECG SHGVCSRGIC QCEEGWVGPT
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                                                                                780
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                                                                                 840
        LDLIQQSQTL FSQHTSRLFY DRIKFLIGKD STHVIPPEVS FDSRRACVIR GQVVAIDGTP
                                                                                900
        LVGVNVSFLH HSDYGFTISR QDGSFDLVAI GGISVILIFD RSPFLPEKRT LWLPWNQFIV
                                                                                960
        VEKVIMORVV SDPPSCDISN PISPNPIVLP SPLTSFGGSC PERGTIVPEL QVVQEEIPIP
                                                                               1020
        SSFVRLSYLS SRTPGYKTLL RILLTHSTIP VGMIKVHLTV AVEGRLTQKW FPAAINLVYT
                                                                               1080
50
        FAWNKTDIYG QKVWGLAEAL VSVGYEYETC PDFILWEQRT VVLQGFEMDA SNLGDWSLNK
        HHILNPQSGI IHKGNGENMF ISQQPPVIST IMGNGHQRSV ACTNCNGPAH NNKLFAPVAL
                                                                               1200
        ASGPDGSVYV GDFNFVRRIF PSGNSVSILE LSTSPAHKYY LAMDPVSESL YLSDTNTRKV
                                                                               1260
        YKLKSLVETK DISKNFEVVA GTGDQCLPFD QSHCGDGGRA SEASLNSERG ITVDRHGFIY
FVDGTMIRKI DENAVITTVI GSNGLTSTQP LSCDSGMDIT QVRLEWFTDL AVNFMDNSLY
                                                                               1320
                                                                               1380
55
        VLDNNIVLQI SENRRVRIIA GRPIHCQVPG IDHFLVSKVA IHSTLESARA ISVSHSGLLF
                                                                               1440
        IAETDERKVN RIQQVTTNGE IYIIAGAPTD CDCKIDPNCD CFSGDGGYAK DAKMKAPSSL
                                                                               1500
        AVSPDGTLYV ADLGNVRIRT ISRNQAHLND MNIYEIASPA DQELYQFTVN GTHLHTLNLI
        TRDYVYNFTY NSEGDLGAIT SSNGNSVHIR RDAGGMPLWL VVPGGQVYWL TISSNGVLKR
                                                                               1620
        VSAQGYNPAL MTYPGNTGLL ATKSNENGWT TVYEYDPEGH LTNATFPTGE VSSFHSDLEK
                                                                               1680
60
        LTKVELDTSN RENVLMSTNL TATSTIYILK QENTQSTYRV NPDGSLRVTF ASGMEIGLSS
                                                                               1740
        EPHILAGAVN PTLCKCNISL PGEHNANLIE WRORKEONKG NVSAFERRLR AHNRNLLSID
                                                                               1800
        FDHITRTCKI YDDHRKFTLR ILYDQTGRPI LWSFVSRYNE VNITYSPSGL VTFIQRGTWN
EKMEYDQSGK IISRTWADGK IWSYTYLEKS VMLLLHSQRR YIFEYDQSDC LLSVTMPSMV
                                                                               1860
                                                                               1920
        RHSLQTMLSV GYYRNIYTPP DSSTSFIQDY SRDGRLLQTL HLGTGRRVLY KYTKQARLSE
                                                                               1980
65
        VLYDTTQVTL TYEESSGVIK TIHLMHDGFI CTIRYRQTGP LIGRQIFRFS EEGLVNARFD
                                                                               2100
        YSYMNFRVTS MQAVINETPL PIDLYRYVDV SGRTEQFGKF SVINYDLNQV ITTTVMKHTK
        IPSANGQVIE VQYEILKAIA YWMTIQYDNV GRHGNMCIRV GVDANITRYF YEYDADGQLQ
                                                                               2160
        TVSVNDKTOW RYSYDLNGDI NLLSHGKSAR LTPLRYDLRD RITRLGEIQY KMDEDGFLRQ
                                                                               2220
        RGNDIFEYNS NGLLQKAYNK ASGWTVQYYY DGLGRRVASK SSLGOHLOFF VDATANPIRV
                                                                               2280
70
        THLYNHTSSE ITSLYYDLOG HLIAMELSSG EEYYVACDNT GTPLAVFSSR GOVIKEILYT
                                                                               2340
        PYGDIYHDTY PDFQVIIGFH GGLYDFLTKL VHLGQRDYDV VAGRWITAYH HIWKQLNLLP
                                                                               2400
        KPFNLYSFEN NYPVGKIQDV AKYTTDIRSW LELFGFQLHN VLPGFPKPEL ENLELTYELL
                                                                               2460
        RLOTKTOEWD PGKTILGIQC ELQKQLRNFI SLDQLPMTPR YNDGRCLEGG KQPRFAAVPS
                                                                               2520
        VFGKGIKFAI KDGIVTADII GVANEDSRRL AAILNNAHYL ENLHFTIEGR DTHYFIKLGS
                                                                               2580
75
        LEEDLVLIGN TGGRRILENG VNVTVSQMTS LLNGRTRRFA DIQLQHGALC FNIRYGTTVE
                                                                               2640
        EEKNHVLEIA RORAVAQAWT KEORRLOEGE EGIRAWTEGE KOOLLSTGRV OGYDGYFVLS
        VEOYLELSDS ANNIHFMROS EIGRR
                                                    SEQ ID NO:231 PFD4 DNA SEQUENCE:
80
        Nucleic Acid Accession #:
                                     NM_000441
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Coding sequence:

225-2567 (underlined sequences correspond to start and stop codons)

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3				1	1	1 .	
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		AATAAAACGT					180
		GGCATTCCGG					240
10		GCCGCCGCAG					300
		GCTCGCTTTC					360
		CCTGGCCAAG					420
		GCCCATCTTG					480
15		TTCGGGAGTT TGCAGTTCCT					540
15		CTTTGGAACA					600 660
		ATCTGTTGTT					720
		TGTATTAAAT					780
••		CAGTGCCCTG					840
20		ATTCATAGTG					900
		CCAAGTGCTG					960
		AGTICTCTCT					1020
		TGCTGATTTC					1080
25		TGATCGGTTT TGCTACTGCC					1140 1200
45		ATCCATCCCA					1260
		GCTGGCTGCA					1320
		AGTATATGCC					1380
20	TTGCCTTTGG	GATCAGCAAC	ATCTTCTCAG	GATTCTTCTC	TTGTTTTGTG	GCCACCACTG	1440
30		CACGGCCGTC					1500
		GATTGTGATG					1560
		CTTGGCAGCT					1620
		TCGTCTGTGG CATCATTCTG					1680 1740
35		GGTCCTGAGA					1800
-		CTACAAAAGT					1860
		ATTTTCCAGT					1920
		AGTTGGATTT					1980
40		GAAACTAATA					2040
40		TTCAACAAAT					2100
		CCCAACCAAG CGTTCCCAAA					2160 2220
		CGTTGTTGGA					2280
		TGTGTATTTT					2340
45		TGACGACAAC					2400
		ACAGAACCAA					2460
		TCAGGATTGT					2520
		TGTCCAGGAT					2580
50		GAGCAAGGAA ACTCATTCTT					2640 2700
30		ATTTATAAAA					2760
		TTTGGCAGCG					2820
		GCTAATAATG					2880
<i></i>		AACAGCCTCT					2940
55		ACTGACCTGG					3000
		TGAAATAAAA					3060
		ATTGGAGTTT GCCGGAATTG					3120 3180
		CCAATATATT					3240
60		ATAAAAATCT					3300
		GACAGATTAT					3360
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		GAGTGAAAAT					3480
65		TTCTCAAAAA					3540
UJ		AAAAAAAATC ACAAAGCATT					3600 3660
		ATACAGGTTT					3720
		GTGCCAACAA					3780
~~		ATATATTTCA					3840
70		GGTCCCCAAT					3900
		GGGATTGGTT					3960
		ACCCCTGCCT					4020
		TGTGTCTATT GGTTGTAAAT					4080 4140
75		ACTTGCTTAT					4200
, ,		GCTCATGCCT					4260
		GATCGAGACC					4320
	AGAAAAAAAG	AAATTAGCCT	AGCGTGGTGG	CTGGCGGGCG	CCTGTAGTCC	CAGCTATTTG	4380
90						GAGCCGAGGT	4440
80	CGTGCCACTG	CACTCCAGCC	TGGGCGACAG	AGCAAGACTC	CGTCTCAAAA	AAAAAAAA	4500

5	AAAGGAAATA GGCTAGAGTT TACTGTCTCT GAAAATTTCA TCCAGTATTG	TGCACTGCTC TGTAAATTCT TCTATGTATT CTTGAAATTA TATATGAGTT	TAATAGTCTT ACTITITIGA GGGTTCATTT TTGTGAATAG AAGCTGCCTT TTAACAAATT AATGTAAAGT	AGGAAATGCC GTGATGACAT TAAGCATAAT TTGTTATATT AAAAAATCAA	AAAGTTACGT AAGTCAGCAA TTTAGTTTTG TTTAACCTAT ATCATGTACA	TTTACAACAA ACTGCGGGAA TATTATCAAT AGGATAAGAT TTTGAAAATA	4560 4620 4680 4740 4800 4860 4920
10	SEQ ID NO:232 Protein Accession	PFD4 Protein secu n #:	<u>ience:</u> <b>O</b> 43511	I			
15							
			21   MVSRPVYSEL				60
20	FFPILTYFIF RDTARVLIAS NVSTKNYNGV PIEVIVTIIA	GTSRHISVGP ALTLLVGIIQ LSIIYTLVEI TAISYGANLE	KEWLLSDVIS FPVVSLMVGS LIFGGLQIGF FQNIGDTNLA KNYNAGIVKS	VVLSMAPDEH IVRYLADPLV DFTAGLLTIV IPRGFLPPEL	FLVSSSNGTV GGFTTAAAFQ VCMAVKELND PPVSLPSEML	LNTTMIDTAA VLVSQLKIVL RFRHKIFVPI AASPSIAVVA	120 180 240 300 360
25	QVAGIISAAI IWVFTCIVSI EPQGVKILRF KNGIISDAVS	VMIAILALGK ILGLDLGLLA SSPIFYGNVD TNNAFEPDED	GNQEFIAFGI LLEPLQKSVL GLIFGLLTVV GFKKCIKSTV IEDLEELDIP KEFQRIDVNV	AAVVIANLKG LRVQFPSWNG GFDAIRVYNK TKEIEIQVDW	MFMQLCDIPR LGSIPSTDIY RLKALRKIQK NSELPVKVNV	LWRQNKIDAV KSTKNYKNIE LIKSGQLRAT PKVPIHSLVL	420 480 540 600 660 720
30			SILETITLIQ				780
35	Nucleic Acid Acc Coding sequence		NM_010 228-100	6029		2 DNA SEQUENCE d to start and stop	
40	1 	11	21 	31 	41 i	51 1	
	GGGCGTGCGC TGCTCCTGCT	GGCCGCAATG CTTGGTGCAG	GACGCGACTC AACTGGGAGC CTGCTGCGCT CGACGCCCAG	TGCTGCTGTG TCCTGAGGGC	GCTGCTGGTG TGACGGCGAC	CTGTGCGCGC CTGACGCTAC	60 120 180 240
45.	TGACTGGAGC TTTCTCTTGT TAGAGAATGG CTGGTTCCCA	CTCGAGTGGA GCTGTCAGCC CAATTTAAAA TGAAGCGGCT	ATTGGTGAGG AGAAGAGTGC GAAAAAGATA ACCAAAGCTG	AGCTGGCTTA ATGAGCTGGA TACTTGTTTT TTCTCCAGGA	CCAGTTGTCT AAGGGTGAAA GCCCCTTGAC GTTTGGTAGA	AAACTAGGAG AGAAGATGCC CTGACCGACA ATCGACATTC	300 360 420 480
50	ACAGAAAGCT CTCACATGAT TATCTGTACC ATGGCCTTCG	AATAGAGCTT CGAGAGGAAG TCTTTCCATT AACAGAACTT	TCCCAGCGTT AACTACTTAG CAAGGAAAGA GGATACTGTG GCCACATACC	GGACGGTGTC TTGTTACTGT CTAGCAAGCA CAGGTATAAT	CTTGACAAAA GAATAGCATC TGCTCTCCGG AGTTTCTAAC	TGTGTTCTGC CTGGGTATCA GGTTTTTTTA ATTTGCCCAG	540 600 660 720 780
55	GCAATAATGG TCAGCATGGC CATATTTGTG AAAGGATTGA	AGACCAGTCC CAATGATTTG GCAATACATG GAACTTTAAG	GTGGAGAATT CACAAGATGA AAAGAAGTTT CCAACCTGGG AGTGGTGTGG	CAACCAGTCG GGATCTCAGA CCTGGTGGAT ATGCAGACTC	TTGTGTGCGG ACAACCTTTC AACCAACAAG TTCTTATTTT	CTGATGTTAA TTGTTAGTAA ATGGGGAAGA AAAATCTTTA	900 960 1020 1080
60	AAACATGAAA ACTTTTTAAT	ACAGCAATCT	TCTTATGCTT TTGCTTCCAA	CTGAATAATC	AAAGACTAAT	GAGAAATGGA TIGTGATIIT AAATAATAAA	1140 1200 1260
65	SEQ ID NO:234   Protein Accession	PFH2 Protein sequ n #:	rence: NP_057	113			
70	GIGEELAYQL ATKAVLQEFG	SKLGVSLVLS RIDILVNNGG	MSQRSLCMDT	KRRCLENGNL SLDVYRKLIE	KEKDILVLPL LNYLGTVSLT	51   MVVWVTGASS DLTDTGSHEA KCVLPHMIER NICPGPVQSN	60 120 180 240
75	IVENSLAGEV	TKTIGNNGDQ		RLMLISMAND		PLLVTYLWQY	300
80	Nucleic Acid Acc	ession #:	NM_000450	SEC	2 ID NO:235 ACC	5 DNA SEQUENCE	•

	Coding sequence	e:	1-1833 (under	lined sequences o	оrrespond to start	and stop codons)	
	1	11 	21	31 .	41	<b>51</b>	
5		CACAGTTTCT					60
		ACAACACCTC ACACACACCT					120
		GCTATTCACC					180 240
10	TGGGTCTGGG	TAGGAACCCA	GAAACCTCTG	ACAGAAGAAG	CCAAGAACTG	GGCTCCAGGT	300
10		ATAGGCAAAA TGTGGAATGA					360 420
		CCAATACATC					480
		AGTGTGACCC					540
15		AATCCCCTGA CTTCCTGCTC					600 660
	ACCATGCAGT	GTATGTCCTC	TGGAGAATGG	AGTGCTCCTA	TTCCAGCCTG	CAATGTGGTT	720
		CTGTGACAAA					780
		GGAACACAAC TTCAGTGTAC					840 900
20	GCTGTGACAT	GCAGGGCCGT	CCGCCAGCCT	CAGAATGGCT	CTGTGAGGTG	CAGCCATTCC	960
		AGTTCACCTT					1020
		CAGCCCAGGT					1080 1140
~~		CTTCTGGCAG					1200
25		TGAAGGGATC					1260
		CATGTGAAGC					1320 1380
		GATTTGAATT					1440
30		AGGTTCCTTC					1500
50		TGAGCTGCAG				AGGACACTGG	1560 1620
						GGTAGCTGGA	1680
		CTGGACTCTC					1740
35		AAAAGCCTTC			GCCAAAGCCT	TGAATCAGAC	1800
40	Protein Accession	ACC5 Protein sequ	NP_000441			•	
	1	11	21	31	44		
4 ==	÷						
	I	Ī	Ĩ	Ĭ	41	51 [	
45		 TLVLLIKESG	 AWSYNTSTEA	 MTYDEASAYC	QQRYTHLVAI	ONKEEIEATN [	60
45	SILSYSPSYY	 TLVLLIKESG WIGIRKVNNV	 AWSYNTSTEA WVWVGTQKPL	 MTYDEASAYC TEEAKNWAPG	QQRYTHLVAI EPNNRQKDED	ONKEEIEYLN CVEIYIKREK	120
45	SILSYSPSYY DVGMWNDERC TALESPEHGS	 TLVLLIKESG WIGIRKVNNV SKKKLALCYT LVCSHPLGNF	AWSYNTSTEA WVWVGTQKPL AACTNTSCSG SYNSSCSISC	 MTYDEASAYC TEEAKNWAPG HGECVETINN DRGYLPSSME	QQRYTHLVAI EPNNRQKDED YTCKCDPGFS TMQCMSSGEW	ONKEEIEYLN CVEIYIKREK GLKCEQIVNC SAPIPACNVV	
	SILSYSPSYY DVGMWNDERC TALESPEHGS ECDAVTNPAN	 TLVLLIKESG WIGIRKVNNV SKKKLALCYT LVCSHPLGNF GFVECFQNFG	AWSYNTSTEA WVWVGTQKPL AACTNTSCSG SYNSSCSISC SPPWNTTCTF	 MTYDEASAYC TEEAKNWAPG HGECVETINN DRGYLPSSME DCEEGFELMG	QQRYTHLVAI EPNNRQKDED YTCKCDPGFS TMQCMSSGEW AQSLQCTSSG	QNKEEIEYLN CVEIYIKREK GLKCEQIVNC SAPIPACNVV NWDNEKPICK	120 180 240 300
45 50	SILSYSPSYY DVGMWNDERC TALESPEHGS ECDAVTNPAN AVTCRAVRQP	TLVLLIKESG WIGIRKVNNV SKKKLALCYT LVCSHPLGNP GFVECFQNPG QNGSVRCSHS	AWSYNTSTEA WWWYGTQKPL AACTNTSCSG SYNSSCSISC SPPWNTTCTF PAGEFTFKSS	 MTYDEASAYC TEEAKNWAPG HGECVETINN DRGYLPSSME DCEEGFELMG CNFTCEEGFM	QQRYTHLVAI EPNNRQKDED YTCKCDPGFS TMQCMSSGEW AQSLQCTSSG LQGPAQVECT	QNKEEIEYLN CVEIYIKREK GLKCEQIVNC SAPIPACNVV NWDNEKPTCK TQGQWTQQIP	120 180 240 300 360
	SILSYSPSYY DVGMWNDERC TALESPEHGS ECDAVTNPAN AVTCRAVRQP VCEAFQCTAL EKPTCEAVRC	TLVLLIKESG WIGIRKVNNV SKKKLALCYT LVCSHPLGNPG GFVECFQNPG QNGSVRCSHS SNPERGYMNC DAVHQPPKGL	AWSYNTSTEA WWWVGTQKPL AACTNTSCSG SYNSSCSISC SPPWNTTCTF PAGEFTFKSS LPSASGSFRY VRCAHSPIGE	 MTYDEASAYC TEEAKNWAPG HGECVETINN DRGYLPSSME DCEEGFELMG CNFTCEEGFM GSSCEFSCEQ FTYKSSCAFS	QQRYTHLVAI EPNNRQKDED YTCKCDPGFS TMQCMSSGEW AQSLQCTSSG LQGPAQVECT GPVLKGSKRL CEEGFELYGS	ONKEEIEYLN CVEIYIKREK GLKCEQIVNC SAPIPACNVV NWDNEKPTCK TQGQWTQQIP QCGPTGEMIN TQLECTSQGQ	120 180 240 300
	SILSYSPSYY DVGMWNDERC TALESPEHGS ECDAVTNPAN AVTCRAVRQP VCEAFQCTAL EKPTCEAVRC WTEEVPSCQV	TLVLLIKESG WIGIRKVNNV SKKKLALCYT LVCSHPLGNF GFVECFQNPG QNGSVRCSHS SNPERGYMNC DAVHQPPKGL VKCSSLAVPG	AWSYNTSTEA WWWVGTQKPL AACTNTSCSG SYNSSCSISC SFPWNTTCTF PAGEFTFKSS LPSASGSFRY WRCAHSPIGE KINMSCSGEP	 MTYDEASAYC TEEAKNWAPG HGECVETINN DRGYLPSSME DCEEGFELMG CNFTCEEGFM GSSCEFSCEQ FTYKSSCAFS VFGTVCKFAC	QQRYTHLVAI EPNNRQKDED YTCKCDPGFS TMQCMSSGEW AQSLQCTSSG LQGPAQVECT GFVLKGSKR CEEGFELYGS PEGWTLNGSA	QNKEEIEYLN CVEIYIKREK GLKCEQIVNC SAPIPACNVV NWDNEKPTCK TQGQWTQQIP QCGPTGEMD TQLECTSQSQ ARTCGATGHW	120 180 240 300 360 420 480 540
	SILSYSPSYY DVGMWNDERC TALESPEHGS ECDAVTNPAN AVTCRAVRQP VCEAFQCTAL EKPTCEAVRC WTEEVPSCQV	TLVLLIKESG WIGIRKVNNV SKKKLALCYT LVCSHPLGNPG GFVECFQNPG QNGSVRCSHS SNPERGYMNC DAVHQPPKGL	AWSYNTSTEA WWWVGTQKPL AACTNTSCSG SYNSSCSISC SFPWNTTCTF PAGEFTFKSS LPSASGSFRY WRCAHSPIGE KINMSCSGEP	 MTYDEASAYC TEEAKNWAPG HGECVETINN DRGYLPSSME DCEEGFELMG CNFTCEEGFM GSSCEFSCEQ FTYKSSCAFS VFGTVCKFAC	QQRYTHLVAI EPNNRQKDED YTCKCDPGFS TMQCMSSGEW AQSLQCTSSG LQGPAQVECT GFVLKGSKR CEEGFELYGS PEGWTLNGSA	QNKEEIEYLN CVEIYIKREK GLKCEQIVNC SAPIPACNVV NWDNEKPTCK TQGQWTQQIP QCGPTGEMD TQLECTSQSQ ARTCGATGHW	120 180 240 300 360 420 480
50	SILSYSPSYY DVGMWNDERC TALESPEHGS ECDAVTNPAN AVTCRAVRQP VCEAFQCTAL EKPTCEAVRC WTEEVPSCQV SGLLPTCEAP	TLVLLIKESG WIGIRKVNNV SKKKLALCYT LVCSHPLGNF GFVECFQNPG QNGSVRCSHS SNPERGYMNC DAVHQPPKGL VKCSSLAVPG	AWSYNTSTEA WWWVGTQKPL AACTNTSCSG SYNSSCSISC SFPWNTTCTF PAGEFTFKSS LPSASGSFRY WRCAHSPIGE KINMSCSGEP	 MTYDEASAYC TEEAKNWAPG HGECVETINN DRGYLPSSME DCEEGFELMG CNFTCEEGFM GSSCEFSCEQ FTYKSSCAFS VFGTVCKFAC LAPFLLWLRK	OQRYTHLVAI EPNNRQKDED YTCKCDPGFS TMQCMSSGEW AQSLQCTSSG LQGPAQVECT GFVLKGSKRL CEEGFELYGS PEGWTLNGSA CLRKAKKFVP	ONKEEIEYIM CVEIYIKREK GLKCEQIVNC SAPIPACNVV MWDNEKPICK TQGQWTQQIP QCGPTGENIN TQLECTSQGQ ARTCGATCHW ASSCQSLESD	120 180 240 300 360 420 480 540 600
50	SILSYSPSYY DVGMWNDERC TALESPEHGS ECDAVTNPAN AVTCRAVRQP VCEAFQCTAL EKPTCEAVRC WTEEVPSCQV SGLLPTCEAP	TLVLLIKESG WIGIRKVNNV SKKKLALCYT LVCSHPLGNF GPVECFQNFG QNGSVRCSHS SNPERGYMNC DAVHQPPKGL VKCSSLAVPG TESNIPLVAG	AWSYNTSTEA WWWVGTQKPL AACTNTSCSG SYNSSCSISC SFPWNTTCTF PAGEFTFKSS LPSASGSFRY WRCAHSPIGE KINMSCSGEP	 MTYDEASAYC TEEAKNWAPG HGECVETINN DRGYLPSSME DCEEGFELMG CNFTCEEGFM GSSCEFSCEQ FTYKSSCAFS VFGTVCKFAC LAPFLLWLRK	OQRYTHLVAI EPNNRQKDED YTCKCDPGFS TMQCMSSGEW AQSLQCTSSG LQGPAQVECT GFVLKGSKRL CEEGFELYGS PEGWTLNGSA CLRKAKKFVP	QNKEEIEYLN CVEIYIKREK GLKCEQIVNC SAPIPACNVV NWDNEKPTCK TQGQWTQQIP QCGPTGEMD TQLECTSQSQ ARTCGATGHW	120 180 240 300 360 420 480 540 600
50 55	SILSYSPSYY DVGMMNDERC TALESPEHGS ECDAVTNPAN AVTCRAVROP VCEAFQCTAL EKPTCEAVRC WTEEVPSCQV SGLLPTCEAP GSYQKPSYIL	TLVLLIKESG WIGIRKVNNV SKKKLALCYT LVCSHPLGNF GFVECFQNPG QNGSVRCSHS SNPERGYMNC DAVHQPPKGL VKCSSLAVPG TESNIPLVAG	AWSYNTSTEA WWWVGTQKPL AACTNTSCSG SYNSSCSISC SPPWNTTCTF PAGEFTFKSS LPSASGSFRY VRCAHSPIGE KINMSCSGEP LSAAGLSLLT N51002	 MTYDEASAYC TEEAKNWAPG HGECVETINN DRGYLPSSME DCEEGFELMG CNFTCEEGFM GSSCEFSCEQ FTYKSSCAFS VFGTVCKFAC LAPFLLWLRK	QORYTHLVAI EPNNRQKDED YTCKCDPGFS TMQCMSSGEW AQSLQCTSSG LQGPAQVECT GFVLKGSKR CEEGFELYGS PEGWTLNGSA CLRKAKKFVP	QNKEEIEYLN CVEIYIKREK GLKCEQIVNC SAPIPACNVV NWDNEKPTCK TQGOWTQQIP QCGPTGEMDN TQLECTSQSQ ARTCGATGHW ASSCQSLESD	120 180 240 300 360 420 480 540 600
50	SILSYSPSYY DVGMMDERC TALESPEHGS ECDAVTNPAN AVTCRAVROP VCEAFQCTAL EKPTCEAVRC WTEEVPSCQV SGLLPTCEAP GSYQKPSYIL Nucleic Acid Acc Coding sequence	TLVLLIKESG WIGIRKVNNV SKKKLALCYT LVCSHPLGNP GPVECFQNPG QNGSVRCSHS SNPERGYMNC DAVHQPPKGL VKCSSLAVPG TESNIPLVAG	AWSYNTSTEA WWWVGTQKPL AACTNTSCSG SYNSSCSISC SPPWNTTCTF PAGEFTFKSS LPSASGSFRY VRCAHSPIGE KINMSCSGEP LSAAGLSLLT N51002 1-3793 (under	MTYDEASAYC TEEAKNWAPG HEGEVETINN DRGYLPSSME DCEEGFELMG CNFTCEEGFM GSSCEFSCEQ FTYKSSCAFS VFGTVCKFAC LAPFLLWLRK SEG	OQRYTHLVAI EPNNRQKDED YTCKCDPGFS TMQCMSSGEW AQSLQCTSSG LQGPAQVECT GFVLKGSKRL CEEGFELYGS PEGWTLNGSA CLRKAKKFVP DID NO:237 PM20	QNKEETEYIM CVETYIKREK GLKCEQTUNC SAPIPACNUV MWDNEKPTCK TQGQWTQQIP QCGPTGEMDN TQLECTSQGQ ARTCGATCHW ASSCQSLESD  B DNA SEQUENCE and stop codons)	120 180 240 300 360 420 480 540 600
50 55	SILSYSPSYY DVGMMNDERC TALESPEHGS ECDAVTNPAN AVTCRAVROP VCEAFQCTAL EKPTCEAVRC WTEEVPSCOV SGLLPTCEAP GSYQKPSYIL Nucleic Acid Acc Coding sequence	TLVLLIKESG WIGIRKVNNV SKKKLALCYT LVCSHPLGNF GFVECFQNFS QNGSVRCSHS SNPERGYMNC DAVHQPPKGL VKCSSLAVPG TESNIPLVAG  ESSION #: 11	AWSYNTSTEA WWWVGTQKPL AACTNTSCSG SYNSSCSISC SPPWNTTCTT PAGEFTFKSS LPSASGSFRY VRCAHSPIGE KINMSCSGEP LSAAGLSLLT  N51002 1-3793 (under	MTYDEASAYC TEEAKNWAPG HGECVETINN DRGYLPSSME DCEEGFELMG CNFTCEEGFM GSSCEFSCEQ FTYKSSCAFS VFGTVCKFAC LAPFLLWLRK  SEG  dined sequences c  31	QQRYTHLVAI QQRYTHLVAI QQRYTHLVAI QCRYCHOPGFS TMQCMSSGEW AQSLQCTSSG LQGPAQVECT GFVLKGSKRL CEEGFELYGS PEGWTLNGSA CLRKAKKFVP DID NO:237 PM2i onespond to start 41	QNKEETEYIM CVETYIKREK GLKCEQIVNC SAPIPACNVV MWDNEKPTCK TQGQWTQQIP QCGPTGEMDN TQLECTSQSQ ARTCGATGHW ASSCQSLESD B DNA SEQUENCE and stop codors) 51	120 180 240 300 360 420 480 540 600
50 55	SILSYSPSYY DVGMMDERC TALESPEHGS ECDAVTNPAN AVTCRAVROP VCEAFQCTAL EKPTCEAVRC WTEEVPSCQV SGLLPTCEAP GSYQKPSYIL  Nucleic Acid Acc Coding sequence  1   ATGATGTGTG	TLVLLIKESG WIGIRVNNV SRKKLALCYT LVCSHPLGNP GPVECFQNPG QNGSVRCSHS SNPERGYMNC DAVHQPPKGL VKCSSLAVPG TESNIPLVAG  ESSION #:  11   AAGTGATGATGCC	AWSYNTSTEA WWWVGTQKPL AACTNTSCSG SYNSSCSISC SFPWNTTCTF PAGEFTFKSS LPSASGSFRY VRCAHSPIGE KINMSCSGEP LSAAGLSLLT  N51002 1-3793 (under 21   CACGATTAAT	MTYDEASAYC TEEAKNWAPG HGECVETINN DRGYLPSSME DCEEGFELMG CNFTCEEGFM GSSCEFSCEQ FTYKSSCAFS VFGTVCKFAC LAPFLLWLRK  SEC  ined sequences c  31   GAGGACACCC	QQRYTHLVAI EPNNRQKDED YTCKCDPGFS TMQCMSSGEW AQSLQCTSSG LQGPAQVECT GFVLKGSKRL CEEGFELYGS PEGWTLNGSA CLRKAKKFVP DID NO:237 PM2: 000espond to start 41   CAATGAGCCA	QNKEETEYIM CVETYIKREK GLKCEQIVNC SAPIPACNVV NMDNEKPTCK TQGQWTQQIP QCGPTGEMIN TQLECTSQGQ ARTCGATCHW ASSCQSLESD  B DNA SEQUENCE and stop codors) 51 AAGGGGGTCC	120 180 240 300 360 420 480 540 600
55	SILSYSPSYY DVGMMNDERC TALESPEHGS ECDAVTNPAN AVTCRAVROP VCEAFQCTAL EKPPCEAVRC WTEEVPSCQV SGLLPTCEAP GSYQKPSYIL  Nucleic Acid Acc Coding sequence  1   ATGATGTGTG CARAGCAGTG	TLVLLIKESG WIGIRKVNNV SKKKLALCYT LVCSHPLGNP GPVECFQNPG QNGSVRCSHS SNPERGYMNC DAVHQPPKGL VKCSSLAVPG TESNIPLVAG  ESION #:  11   AAGTGATGCC GCTCGGACTC	AWSYNTSTEA WWWVGTQKPL AACTNTSCSG SYNSSCSISC SPPWNTTCTF PAGEFTFKSS LPSASGSFRY VRCAHSPIGE KINMSCSGEP LSAAGLSLLT  N51002 1-3793 (under 21   CACGATTAAT AGACTCCCAT	MTYDEASAYC TEEAKNWAPG HGECVETINN DRGYLPSSME DCEEGFELMG CNFTCEEGFM GSSCEFSCEQ FTYKSSCAFS VFGTVCKFAC LAPFLLWLRK SEG ined sequences c 31   GAGGACACCC TTTGAGCAGC	OQRYTHLVAI EPNNRQKDED YTCKCDPGFS TMQCMSSGEW AQSLQCTSSG LQGPAQVECT GFVLKGSKRL CEEGFELYGS PEGWTLNGSA CLRKAKKFVP DID NO:237 PM2I OMESPOND TO STATE  CAATGAGCCA TGATGAGCCA TGATGATGA	ONKEEIEYLM CVEIYIKREK GLKCEQIVNC SAPIPACNVV MWDNEKPICK TQGQWTQQIP QCGPTGENDN TQLECTSQGQ ARTCGATCHW ASSCQSLESD  B DNA SEQUENCE and stop codors)  51   AAGGGGGTCC TATGCTAGAT	120 180 240 300 420 480 540 600
50 55	SILSYSPSYY DVGMMNDERC TALESPEHGS ECDAVTNPAN AVTCRAVROP VCEAFQCTAL EKPTCEAVRC WTEEVPSCOV SGLLPTCEAP GSYQKPSYIL  Nucleic Acid Acc Coding sequence  1   ATGATGTGTG CAAAGCAGTG GAAAGGGATC CAAAGACTTC	TLVLLIKESG WIGIRKVNNV SKKKLALCYT LVCSHPLGNF GFVECFQNFS GNGSVRCSHS SNPERGYMNC DAVHQPPKGL VKCSLAVPG TESNIPLVAG  ESSION #:  11   AAGTGATGCC GCTCGGACTC GTCTTCTAGA AGGATGTCAT	AWSYNTSTEA WWWVGTQKPL AACTNTSCSG SYNSSCSISC SPPWNTTCTT PAGEFTFKSS LPSASGSFRY VRCAHSPIGE KINMSCSGEP LSAAGLSLLT  N51002 1-3793 (under 21   CACGATTAAT AGACTCCAT CACCTTCGG CTATGACCGA	MTYDEASAYC TEEAKNWAPG HGECVETINN DRGYLPSSME DCEEGFELMG CNFTCEEGFM GSSCEFSCEQ FTYKSSCAFS VFGTVCKFAC LAPFLLWLRK  SEG dined sequences c  31   GAGGACACCC TTTGAGCAGC GAGACCCAGG GAGACCCAGG GAGACCCACGG GAGACCCACGG GAGACCCACGC	QQRYTHLVAI QQRYTHLVAI QQRYTHLVAI QQRYTHLVAI QUENTIAN QUEN	QNKEETEYLM CVETYIKREK GLKCEQTVNC SAPIPACNVV MWDNEKPTCK TQGQWTQQIP QCGPTGEMDN TQLECTSQSQ ARTCGATGHW ASSCQSLESD  B DNA SEQUENCE and stop codons) 51 AAGGGGGTCC TATGCTAGAT ACTTGCCCAG CAATTCAGCC	120 180 240 300 360 420 480 540 600
55	SILSYSPSYY DVGMMNDERC TALESPEHGS ECDAVTNPAN AVTCRAVROP VCEAFQCTAL EKPTCEAVRC WTEEVPSCOV SGLLPTCEAP GSYQKPSYIL  Nucleic Acid Acc Coding sequence  1   ATGATGTGTG GAAAGCAGTG GAAAGGGATC CAAAGACTTC CTGCCACAGG	TLVLLIKESG WIGIRKVNNV SKKKLALCYT LVCSHPLGNF GFVECFQNFG QNGSVRCSHS SNPERGYMNC DAVHQPPKGL VKCSSLAVPG TESNIPLVAG  22 23 21 1 AAGTGATGCC GCTCGGACTC GTCTTCTAGA AGGATGTCAT ATATCGAATC	AWSYNTSTEA WWWVGTQKPL AACTNTSCSG SYNSSCSISC SFPWNTTCTF PAGEFTFKSS LPSASGSFRY VRCAHSPIGE KINMSCSGEP LSAAGLSLLT  N51002 1-3793 (under 21   CACGATTAAT AGACTCCAT CACCCTTCGG CTATGACCGA CCTAACAGGA	MTYDEASAYC TEEAKNWAPG HGECVETINN DRGYLPSSME DCEEGFELMG CNFTCEEGFM GSSCEFSCEQ FTYKSSCAFS VFGTVCKFAC LAPFLLWLRK  SEG  ined sequences c  31   GAGGACACCC TTTGAGCAGC GAGCACAGG GACTCACTCC GGGCTGCTG	QQRYTHLVAI EPNNRQKDED YTCKCDPGFS TMQCMSSGEW AQSLQCTSSG LQGPAQVECT GFVLKGSKRL CEEGFELYGS PEGWTLNGSA CLRKAKKFVP  ID NO:237 PM2i omespond to start 41   CAATGAGCCCAT TGATGGTGAA AAGCCTCTA AAGGCATGTTAAGGG GTTCTAAGGG GTTCTAAGGG	QNKEEIEYLM CVEIYIKREK GLKCEQIVNC SAPIPACNVV NWDNEKPICK TQGQWTQQIP QCGPTGEMIN TQLECTSQGQ ARTCGATCHW ASSCQSLESD  B DNA SEQUENCE and slop codons) 51 AAGGGGGTCC TATGCTAGAT ACTTGCCCAG GCATTCAACCC GGCTGATCCA	120 180 240 300 360 420 480 540 600
50 55 60 65	SILSYSPSYY DVCMMNDERC TALESPEHGS ECDAVTNPAN AVTCRAVROP VCEAPQCTAL EKPPCEAVRC WTEEVPSCQV SGLLPTCEAP GSYQKPSYIL  Nucleic Acid Acc Coding sequence  1   ATGATGTGTG GAAAGCAGTG GAAAGCAGTC CAAAGACTTC CTGCCACAGG CCGGAATTTG	TLVLLIKESG WIGIRKVNNV SKKKLALCYT LVCSHPLGNF GFVECFQNFS GNGSVRCSHS SNPERGYMNC DAVHQPPKGL VKCSLAVPG TESNIPLVAG  ESSION #:  11   AAGTGATGCC GCTCGGACTC GTCTTCTAGA AGGATGTCAT	AWSYNTSTEA WWWVGTQKPL AACTNTSCSG SYNSSCSISC SPPWNTTCTF PAGEFTFKSS LPSASGSFRY VRCAHSPIGE KINMSCSGEP LSAAGLSLLT  N51002 1-3793 (under 21 i CACGATTAAT AGACTCCCAT CACCTTCGG CTATGACCGA ACTACAGGA AAAAGAATTA	MTYDEASAYC TEEAKNWAPG HEBCVETINN DRGYLPSSME DCEEGFELMG GSSCEFSCEQ FTYKSSCAFS VFGTVCKFAC LAPFLLWLRK  SEG  ined sequences c  31   GAGGACACCC TTTGAGCAGC GAGACCCAGG GACTCACTCC GGGCTGGCTG AATGCCTGCA	OQRYTHLVAI EPNNRQKDED YTCKCDPGFS TMQCMSSGEW AQSLQCTSSG EQPAQVECT GFVLKGSKRL CEEGFELYGS PEGWTLNGSA CLRKAKKFVP  ID NO:237 PM2 OMESPOND TO STATE CAATGAGCCA TGATGGTGAA AAAGCCTCTC GTTCTAAAGGG GGGAACAACT	ONKEETEYLM CVETYIKREK GLKCEQIVNC SAPIPACNVV MWDNEKPTCK TQGQWTQQIP QCGPTGEMIN TQLECTSQGQ ARTCGATCHW ASSCQSLESD  B DNA SEQUENCE and stop codons) 51   AAGGGGGTCC TATGCTAGAT ACTTGCCCAG CACTTCAGCCA TCTAGAAAAG	120 180 240 300 360 420 480 540 600
55	SILSYSPSYY DVGMMNDERC TALESPEHGS ECDAVTNPAN AVTCRAVROP VCEAFQCTAL EKPTCEAVRC WTEEVPSCOV SGLLPTCEAP GSYQKPSYIL  Nuclec Acid Acc Coding sequence  1   ATGATGTGTG CAAAGCAGTG GAAAGCAGTC CAAAGCATTC CTGCCACAGG CCGGAATTTC GAAAGAAAA TTGGAGTGCC	TLVLLIKESG WIGIRKVNNV SKKKLALCYT LVCSHPLGNF GFVECFONFG QNGSVRCSHS SNPERGYMNC DAVHQPPKGL VKCSSLAVPG TESNIPLVAG  #: 11   AAGTGATGCC GCTCGGACTC GTCTTCTAGA AGGATGTCAT ATATCGAATT CTGCACTGACT TTCTGTGACT	AWSYNTSTEA WWWVGTQKPL AACTNTSCSG SYNSSCSISC SFPWNTTCTF PAGEFTFKSS LPSASGSFRY VRCAHSPIGE KINMSCSGEP LSAAGLSLLT  N51002 1-3793 (under 21   CACCATTAAT AGACTCCAT CACCCTTCGG CTATGACCGA AAAAGAATTA ACACTGAAAGA ACATGAAAGA ACATGAAAGA	MTYDEASAYC TEEAKNWAPG HGECVETINN DRGYLPSSME DCEEGFELMG GSSCEFSCEQ FTYKSSCAFS VFGTVCKFAC LAPFLLWLRK  SEC  ined sequences c  31   GAGGACACCC TTTGAGCAGC GAGACCCAGG GAGACCCAGG GAGACCACTC GGGCTGGCTG AATGCCTGCA ACCTAAGAA	QQRYTHLVAI QQRYTHLVAI QQRYTHLVAI QQRYTHLVAI QUENTE CEPNING COMPA QSLQCTSSG LQGPAQVECT GFVLKGSKRL CEEGFELYGS PEGWTLINGSA CLRKAKKFVP  ID NO:237 PM2I OMESPOND TO STATE  41   CAATGAGCCA TGATGGTGAA AAAGCCTCTC AGAGACAGCT GTTCTAAGGG GGGAACAACT TGACGGTGGT TGACGGTGGT TGACGGTGGT	QNKEETEYLM CVETYIKREK GLKCEQTVNC SAPIPACNVV NWDNEKPTCK TQGQWTQQIP QCGPTGEMDN TQLECTSQGQ ARTCGATGHW ASSCQSLESD  B DNA SEQUENCE and stop codons) 51 AAGGGGGTCC TATGCTAGAT ACTTGCCCAG CAATTCAGCC GGCTGATCCA TCTAGAAAAG ACTGGAGCAT AAAACGGCAA	120 180 240 300 360 420 480 540 600 120 180 240 360 420 480
50 55 60 65	SILSYSPSYY DVCMMNDERC TALESPENGS ECDAVTNPAN AVTCRAVROP VCEAFQCTAL EKPPCEAVRC WTEEVPSCQV SGLLPTCEAP GSYQKPSYIL  Nucleic Acid Acc Coding sequence  1   ATGATGTGTG GAAAGGATTC CAAAGACTTC CCAAAGACTTC CAAAGAATTG GAAAGAATTG GAAAGAATTG GAAAGAATTG GAAGAAGAATTG GAAGAAGAATTG GAAGAAGAATTG GAAGAAGAATTG GAAGAAGAATTG GAAGAAGAAC ATTGGAAGTGCC GCCCAGTCTC	TLVLLIKESG WIGIRKVNNV SKKKLALCYT LVCSHPLGNF GFVECFQNPG QNGSVRCSHS SNPERGYMNC DAVHQPPKGL VKCSSLAVPG TESNIPLVAG  22 23 21 1 AAGTGATGCC GCTCGGACTC GTCTTCTAGA AGGATGTCAT ATATCGAATC CTGCACTGAC TCTCTGAACT TCTCTGAACT TCTCTGAACT TCTCTGAACT CTCTCAACG CCTCAGGAGT	AWSYNTSTEA WWWVGTQKPL AACTINTSCSG SYNSSCSISC SFPWNTTCTF PAGEFTFKSS LPSASGSFRY VRCAHSPIGE KINMSCSGEP LSAAGLSLLT  N51002 1-3793 (under 21   CACGATTAAT AGACTCCAT CACCCTTCGG CCTATCAGGA AAAGAATTA TAAAGCTGAA ACATGAAAGA ATCCAGTGAA	MTYDEASAYC TEEAKNWAPG HGECVETINN DRGYLPSSME DCEEGFELMG GSSCEFSCEQ FTYKSSCAFS VFGTVCKFAC LAPFLLWLRK  SEG  ined sequences c  31   GAGGACACCC TTTGAGCAGC GAGTACCCC GGGCTGCTG AATGCCTGCA AGAACAACA TCACTAAGAA GTTGAAGTTC	QQRYTHLVAI EPNNRQKDED YTCKCDPGFS TMQCMSSGEW AQSLQCTSSG LQGPAQVECT GFVLKGSKRL CEEGFELYGS PEGWTLNGSA CLRKAKKFVP  ID NO:237 PM2i omespond to start 41   CAATGAGCCAT TGATGGTGAA AAAGCCTTCT GTTCTAAGGG GGGAACAACT CAAGGACTACT TGACGGTGGT TCAAGGGCACT TCAAGGCACT TCAAGGCACT TCAAGGCACT TCAAGGCACT TCAAGGCACT TCAAGGCACT TCAAGGCACT TCAAGGCACT TCAAGGCACT	QNKEETEYLM CVETYIKREK GLKCEQTVNC SAPIPACNVV NMDNEKPTCK TQGQWTQQIP QCGPTGEMIN TQLECTSQGQ ARTCGATCHW ASSCQSLESD  B DNA SEQUENCE and slop codons)  51 AAGGGGGTCC TATGCTAGAT ACTTGCCCAG GCTGATCCA TCTAGAAAAG ACTGGAGCAT TCTAGAAAAG AAAACGCAA GAAATCTTTG	120 180 240 300 360 420 480 540 600 120 180 240 300 360 420 480 540
50 55 60 65	SILSYSPSYY DVCMMNDERC TALESPEHGS ECDAVTNPAN AVTCRAVROP VCEAFQCTAL EKPPCEAVRC WTEEVPSCQV SGLLPTCEAP GSYQKPSYIL  Nucleic Acid Acc Coding sequence T ATCATGTOTG GAAAGCAGTG GAAAGGGATC CTGCCACAGG CCGGAATTTG GAAGAAGAAA TTGGAGTGCC TTTGAGCACC	TLVLLIKESG WIGIRKVNNV SKKKLALCYT LVCSHPLGNF GFVECFONFG QNGSVRCSHS SNPERGYMNC DAVHQPPKGL VKCSSLAVPG TESNIPLVAG  #: 11   AAGTGATGCC GCTCGGACTC GTCTTCTAGA AGGATGTCAT ATATCGAATT CTGCACTGACT TTCTGTGACT	AWSYNTSTEA WWWVGTQKPL AACTNTSCSG SYNSSCSISC SPPWNTTCTF PAGEFTFKSS LPSASGSFRY VRCAHSPIGE KINMSCSGEP LSAAGLSLLT  N51002 1-3793 (under 21   CACGATTAAT AGACTCCAT CACCTTCGG CTATGACCGA ACACGGAA ACACGGAA ACACGGAA ACACGGAA ACACGGAA ACACGGAA ACACGGAA ACACGGAAAAGGAATAA GGATGAAAAG GGATGAAAAG	MTYDEASAYC TEEAKNWAPG HEBCVETINN DRGYLPSSME DCEEGFELMG CNFTCEEGFM GSSCEFSCEQ FTYKSSCAFS VFGTVCKFAC LAPFLLWLRK  SEG  ined sequences c  31   GAGGACACCC TTTGAGCAGC GAGTCCACTGC GGGCTGGCTG AATGCCTGCA AGAACAACA CCACTAAGAA CTTCAAGCAGC GTTAAGGGAGC GTTAAGGGAGC GTTAAGGGAGC GTTAAGGGAGC	OQRYTHLVAI EPNNRQKDED YTCKCDPGFS TMQCMSSGEW AQSLQCTSSG EQSLQCTSSG CEGFELYGS PEGWTLNGSA CLRKAKKFVP  ID NO:237 PM2  CRATGAGCCA TGATGAGCCA TGATGAGCCA TGATGAGCCA TGTCTAAGGG GGGAACAACT CAAGACTATT TGACGGTGGT TCAAGGCACT TGAAGGCACT TGAAGGCACT TGAAGGCACT TGAAGGCACT CAAGACTATT TCAAGGCACT GACTGAGGGT	ONKEEIEYLM CVEIYIKREK GLKCEQIVNC SAPIPACNVV MWDNEKPICK TQGQWTQQIP QCGPTGENDN TQLECTSQGQ ARTCGATCHW ASSCQSLESD  B DNA SEQUENCE and stop codons)  51  AAGGGGGTCC TATGCTAGAT ACTTGCCCAG CAATTCAGCC GGCTGATCCA TCTAGAAAAG ACTGGACCAT AAAACGCCAA AAAACGCCAA AAAACGCCAA AGAATCTTG TTCTTTAGAA	120 180 240 300 420 480 600 120 180 240 420 480 540 600
50 55 60 65	SILSYSPSYY DVGMMNDERC TALESPEHGS ECDAVTNPAN AVTCRAVROP VCEAFQCTAL EKPTCEAVRC WTEEVPSCOV SGLLPTCEAP GSYQKPSYIL  Nucleic Acid Acc Coding sequence  1   ATGATGTGTG GAAAGCAGTG CAAAGCAGTC CAAAGACTTC CTGCCACAGG CCGGAATTTG GAAGAAGAAA TTGGAGTGCC GCCCAGTCTC TTTGAGCACC GAACAAAATG GAACAAAATG	TLVLLIKESG WIGIRKVNNV SKKKLALCYT LVCSHPLGNF GFVECFQNFG QNGSVRCSHS SNPERGYMNC DAVHQPPKGL VKCSSLAVPG TESNIPLVAG  ESSION #:  11	AWSYNTSTEA WWWVGTQKPL AACTNTSCSG SYNSSCSISC SFPWNTTCTF PAGEFTFKSS LPSASGSFRY VRCAHSPIGE KINMSCSGEP LSAAGLSLLT  N51002 1-3793 (under 21   CACCATTAAT AGACTCCAT CACCTTCGG CCTATGACCGA CCTACCAGGA AAAAGAATTA ACATGAAAGA ACATGAAAGA ATCCAGTGAA ACATGAAAGA ATCCAGTGAA ACATGAAAAAATG AGAACTACCT AAGAAAAATTC AAGAAAAATTC AAGAAAAAATTC AAGAAAAAATTC AAGAAAAAATTC	MTYDEASAYC TEEAKNWAPG HGECVETINN DRGYLPSSME DCEEGFELMG CNFTCEEGFM GSSCEFSCEQ FTYKSSCAFS VFGTVCKFAC LAPFLLWLRK  SEC  ined sequences c  31   GAGGACACCC TTTGAGCAGC GAGACCCAGG GACTCACTCC GGGCTGGCTG AATGCCTGCA ACAACAA TCACTAAGAA GTTGAAGGTC GCTGCTAATC GCATCAAGCG GCATCAAGCG GCATCAAGCG GCATCAAGCG	QQRYTHLVAI QQRYTHLVAI QQRYTHLVAI QQRYTHLVAI QUENTICKEDPGFS TMQCMSSGEW AQSLQCTSSG LQGPAQVECT GFVLKGSKRL CEEGFELYGS PEGWTLNGSA CLRKAKKFVP  ID NO:237 PM2I OMESPOND TO STAN AAAGCCTCT AGAGACAGCT GATTAAGGG GGGAACAACT CAAGGCTATT TGACGGTGGT TCAAGGCTT TCAAGGCACT TCAAGGCACT AGGACTTAT TGACGGTGGT TCAAGGCACT AGGACTTAT TGACGGTTGT AGGAGATTGT AGGAGATTGT AGGAGATTGT AGGAGATTGT AGGAGATTGT AGGAGATTGA	ONKEETEYIM CVETYIKREK GLKCEQIVNC SAPIPACNVV NWDNEKPTCK TQGQWTQQIP QCGPTGEMDN TQLECTSQGQ ARTCGATCHW ASSCQSLESD  B DNA SEQUENCE and slop codons) 51 AAGGGGGTCC TATGCTAGAT ACTTGCCCAG CAATTCAGCC GGCTGATCCA TCTAGAAAAG ACTGGAGCAT AAAACGCAA GAAATCTTTG TTCTTTAGAA TGCCTTGCGT AGGAGTAAAAAG GAATCTTTG TCTTTTTAGAA TGCCTTCGGT AGAGTCAGAA	120 180 240 360 420 480 540 600 120 180 240 360 420 480 540 660 720
50 55 60 65	SILSYSPSYY DVCMMNDERC TALESPEHGS ECDAVTNPAN AVTCRAVROP VCEAPQCTAL EKPPCEAVRC WTEEVPSCQV SGLLPTCEAP GSYQKPSYIL  Nucleic Acid Acc Coding sequence  1   ATGATGTGTG GAAAGCAGTG CAAAGCAGTTC CTACCACAGG CCGGAATTTG GAAGAAGACTCTC TTTGAGCACC AGAGTCTCTC TTTGAGCACC AGAGTCTCTC TTTGAGCACC AGAGTCTCTG GAACAAAATG CATCTTGAAG CATCTTTGAAG CATCTTTGAAGC CATCTTTGAAGC CATCTTTGAAGC CATCTTTGAAGC CATCTTTGAAGC CATCTTTGAAGC CATCTTTGAAGC CATCTTTGAAGC CATCTTTGAAGC CATCTTTGAAG	TLVLLIKESG WIGIRKVNNV SKKKLALCYT LVCSHPLGNF GFVECFQNPG QNGSVRCSHS SNPERGYMNC DAVHQPPKGL VKCSSLAVPG TESNIPLVAG  22 22 23 24 24 25 25 26 27 27 27 27 27 27 27 27 27 27 27 27 27	AWSYNTSTEA WWWYGTQKPL AACTINTSCSG SYNSSCSISC SFPWNTTCTF PAGEFTFKSS LPSASGSFRY VRCAHSPIGE KINMSCSGEP LSAAGLSLLT  N51002 1-3793 (under 1-3793 (under CACCTTCGG CCTATGACGGA AAAGATTAAT AGACTCCAT TAAAGCTGAA ACATGAAGA ATCCAGTGAA GGATGAAAG GGATGAAAG GGATGAAAG GGATGAAAG TGGACAGAAA	MTYDEASAYC TEEAKNWAPG HGECVETINN DRGYLPSSME DCEEGFELMG GSSCEFSCEQ FTYKSSCAFS VFGTVCKFAC LAPFLLWLRK  SEG  ined sequences c  31   GAGGACACCC TTTGAGCAGC GAGTCACTC GGGCTGCTG AATGCCTGCA AGAACAACA TCACTAAGAA GTTGAAGTTC GTAAGGAGC GCTGCTAATC GCATCAAGC GCATCAACC GCATCAAGC GCATCAAGC GCATCAAGC GCATCAAGC GCATCAAGC GCATCAAGC GCATCAAGC GCATCAAGC GCATCAAGC GTCCATGAGA	QQRYTHLVAI   QQRYTHLVAI   EPNNRQKDED   YTCKCDPGFS   TMQCMSSGEW    AQSLQCTSSG    CQSPAQVECT    GEVILKGSKRL    CEEGFELYGS    PEGWTLNGSA    CLRKAKKFVP    ID NO:237 PM2    Omespond to start   CAATGAGCCA    CAATGAGCCA    CAATGAGCCA    GTTCTAAGGG    GGGAACAACT    CAAGGCATCATC    TGACGGTGTT   TGACGGTGTT   TGACGGTGTT   TCAAGGCACT    AGGGATTCTAAGGGTACCACAGGGATTCTC    AGGGATTCTCAAGGGTTCTCCAAGCGTTTTTCTC	ONKEETEYLM CVETYIKREK GLKCEQIVNC SAPIPACNVV NMDNEKPTCK TQGQWTQQIP QCGPTGEMIN TQLECTSQGQ ARTCGATCHW ASSCQSLESD  B DNA SEQUENCE and slop codons)  51 AAGGGGGTCC TATGCTAGAT ACTTGCCCAG GCTGATCCA TCTAGAAAAG ACTGGACCAT TCTAGAAAAG ACTGGACCAT TCTAGAAAAG GAATCTTTG TTCTTTAGAA TGCCTTGCGT AGAGTAGAAA CAATGGTTCT	120 180 240 300 360 420 540 600 120 180 240 360 420 420 600 600 600 600 720 780
50 55 60 65	SILSYSPSYY DVCMMNDERC TALESPEHGS ECDAVTNPAN AVTCRAVROP VCEAFQCTAL EKPPCEAVRC WTEEVPSCQV SGLLPTCEAP GSYQKPSYIL  Nucleic Acid Acc Coding sequence TATEGATGTGTG GAAAGGGATC CAAAGACTTC GACAGAGACTTG GACAGAGACTC TTTGAGCACC AGACTACTC GAACAAAATG CATTGAAG ATAGACTCAA ATAGACTCAA	TLVLLIKESG WIGIRKVNNV SKKKLALCYT LVCSHPLGNP GPVECFQNPG QNGSVRCSHS SNPERGYMNC DAVHQPPKGL VKCSSLAVPG TESNIPLVAG  25	AWSYNTSTEA WWWYGTQKPL AACTNTSCSG SYNSSCSISC SPPWNTTCTF PAGEFTFKSS LPSASGSFRY VRCAHSPIGE KINMSCSGEP LSAAGLSLLT  N51002 1-3793 (under 21   CACGATTAAT AGACTCCCAT CACCCTTCGG CTATGACCGA AAAAGAATTA ACATGAAAA ACATGAAAA ACATGAAAAG GGATGAAAAG GGATGAAAAG GGATGAAAAG TGGACAGAAA TTGGACAGAAA	MTYDEASAYC TEEAKNWAPG HEGEVETINN DRGYLPSSME DCEEGFELMG GSSCEFSCEQ FTYKSSCAFS VFGTVCKFAC LAPFLLWLRK  SEG  ined sequences c  31   GAGGACACCC TTTGAGCAGC GAGACCAGG GACTCACTCC GGGCTGGCTG AATGCCTGCA AGAACAACA TCACTAAGAA TCACTAAGAA GTTGAAGTAC GTAAGGGAGC GCTATCACG GTCATCACG GTCATCACG GTCATCACG GTCATCACG GTCATCACG GTTGAACTAC GTTGAACTAC GTTGAACTAC	OQRYTHLVAI EPNNRQKDED YTCKCDPGFS TMQCMSSGEW AQSLQCTSSG EQSLQCTSSG CLGPAQVECT GFVLKGSKRL CEEGFELYGS PEGWTLNGSA CLRKAKKFVP  ONESPOND TO SLAN  L CAATGAGCCA TGATGGTGAA AAAGCCTCTC AGAGCAACT CTCAAGGG GGGAACAACT TGACGGTGGT TCAAGGG GGGAACAACT TCAAGGGATCAT TCAAGGATTGT AGGGATCTGT AGGGATCTGT AGGGATCTGT AGGGATCTGT AGGGATCTGTC AAGAATTGCT AAGAATTGCT AAGAATTGCT	ONKEEIEYLM CVEIYIKREK GLKCEQIVNC SAPIPACNVV MWDNEKPICK TQGQWTQQIP QCGPTGEMIN TQLECTSQGQ ARTCGATCHW ASSCQSLESD  B DNA SEQUENCE and stop codons)  51  AAGGGGGTCC TATGCTAGAT ACTTGCCCAG CCATTCAGCC TCTAGAAAAG ACTGAGCAT AAAACGCAA AAAACGTAGAT TCTTTTAGAA TGCCTTGCGT AGAGTCAGAA CAATGGTTCT TGAAAAGCAA	120 180 240 360 420 480 600 120 180 240 360 420 480 540 660 720 780 840
50 55 60 65	SILSYSPSYY DVGMMNDERC TALESPEHGS ECDAVTNPAN AVTCRAVROP VCEAFQCTAL EKPTCEAVRC WTEEVPSCOV SGLLPTCEAP GSYQKPSYIL  Nucleic Acid Acc Coding sequence  1   ATGATGTGTG GAAAGCAGTG CAAAGCAGTC CAAAGACTTC CTGCCACAGG CCGGAATTTG CAAGAGTATTG GAAGAAGAAA TTGGAGTACC TTTGAGCACC GCCCAGTCTC GAACAAAATG CATCTTGAAG ATAGACTCTG GAACAAAATG CATCTTGAAGA ATAGACTCAA ATGACTCAAA ATGACTCAAA GTGGAACAGG	TLVLLIKESG WIGIRKVNNV SKKKLALCYT LVCSHPLGNF GFVECFQNFG QNGSVRCSHS SNPERGYMNC DAVHQPPKGL VKCSSLAVFG TESNIPLVAG  ###################################	AWSYNTSTEA WWWYGTQKPL AACTINTSCSG SYNSSCSISC SFPWNTTCTF PAGEFTFKSS LPSASGSFRY VRCAHSPIGE KINMSCSGEP LSAAGLSLLT  N51002 1-3793 (under 21   CACCATTAAT AGACTCCAT CACCTTCGG CTATGACCGA CCTACGAGA AAAAGAATTA ACATGAAAGA ACATGAAAGA ACATGAAAGA ATCCAGTGAA ACATGAAAGA ATCCAGTGAA ACATGAAAGA ATCCAGTGAA ACATGAAAAGA TAGAACTACCT AAGAAAAATT TGGACAGAAA TAGTCAAATA CGAAGAAAAT AGGACAGAAA	MTYDEASAYC TEEAKNWAPG HGECVETINN DRGYLPSSME DCEEGFELMG GSSCEFSCEQ FTYKSSCAFS VFGTVCKFAC LAPFLLWLRK  SEG  ined sequences c  31   GAGGACACCC TTTGAGCAGC GAGACCCAGG GACTCACTCC GGGCTGGCTG AATGCCTGCA TCACTAAGAA GTTGAAGTAC GCTGCTAATC GCATCAAGCA GTTGAACTAC GTTGAACTAC GTTGAACTAC GTTGAACTAC GTTGAACTAC GTTGAACTAC GTTGAACTAC GTTGAACTAC TTAGCAGCC GTTCTATCATTA	QQRYTHLVAI QQRYTHLVAI QQRYTHLVAI PRINEQKDED YTCKCDPGFS TMQCMSSGEW AQSLQCTSSG CLQCTSSG LQGPAQVECT GFVLKGSKRL CEEGFELYGS PEGWTLNGSA CLRKAKKFVP  ID NO:237 PM2I OMESPOND TO STAN AAAGCCTCTC AGAGACAGCT CAAGAGCTATC CAAGAGCTATT TGACGGTGGT TCAAGGGTT TCAAGGGTT CAAGGATTCAC AGGAATTGCT AGGGATTCAC AGGAATTGCT AGGGATTCAC AGGAATTGCT TTTCTTCCCG AAACAGAAGA	ONKEETEYIM CVETYIKREK GLKCEQTVNC SAPIPACNVV NWDNEKPTCK TQGQWTQQIP QCGPTGEMDN TQLECTSQGQ ARTCGATCHW ASSCQSLESD  B DNA SEQUENCE and slop codons) 51 ARGGGGGTCC TATGCTAGAT ACTTGCCAG CAATTCAGCC GGCTGATCCA TCTAGAAAAG ACTGGAGCAT AAAACGCAA GAAATCTTTG TTCTTTAGAA TGCCTTCCGT AGGGTCAGAA CAATGGTTCT TGAAAACCAA AATGGACAA CAATGGTCAGAA CAATGGTCAGAA CAATGGTCAGAA CAATGGTCAGAA CAATGGTCAGAA CAATGGTCT TGAAAACCAC AGTGGAGAGA AATGAACACC	120 180 240 360 420 480 540 600 120 180 240 360 420 480 540 660 720 780 840 960
50 55 60 65	SILSYSPSYY DVCMMNDERC TALESPEHGS ECDAVTNPAN AVTCRAVROP VCEAPQCTAL EKPPCEAVRC WTEEVPSCOV SGLLPTCEAP GSYQKPSYIL  Nucleic Acid Acc Coding sequence  1   ATGATGTGTG GAAAGCAGTG GAAAGACTTC CTACCACAGG CCGGAATTTG GAAGAACTC TTTGAGCACC ACAGTCTCT TTTGAGCACC ACAGTCTCT CTTTGAGCACC ACAGTCTTCT CATCTGAGAAATG CATCTTTGAAG ATGATTATGAA ACTTATGAAA ACTTATGAAA ACTTATGAAA ACTTATGAAA ACTTATGAAA ACTTATGAAA ACTTATGAAAA ACTTATGAAAAAAACAGAAAAAAAAAA	TLVLLIKESG WIGIRKVNNV SKKKLALCYT LVCSHPLGNF GFVECFONFE QNGSVRCSHS SNPERGYMNC DAVHOPPKGL VKCSSLAVPG TESNIPLVAG    ESSION #:  11	AWSYNTSTEA WWWYGTQKPL AACTINTSCSG SYNSSCSISC SFPWINTTCTF PAGEFTFKSS LPSASGSFRY VRCAHSPIGE KINMSCSGEP LSAAGLSLLT  N51002 1-3793 (under 1-3793 (under CACCTTCGG CTATGACGGA AAAGATTAAT AGACTCCAT TAAAGCTGAA ACATGAAGA ATCCAGTGAA GGATGAAAG GGATGAAAG GGATGAAAG GGATGAAAG AGAAAAATT TGGACAGAAA TAGTCAAATA GAAAGAACT TGGACAGAAA TAGTCAAATA GAAAGAACGT TGGACAGAAA TAGTCAAATA GAAAGAACGT TGGACAGAAA TAGTCAAATA GAAAGAACGT TGGACAGAAA GGAAGAAAG GGAGGCCATG	MTYDEASAYC TEEAKNWAPG HGECVETINN DRGYLPSSME DCEEGFELMG GSSCEFSCEQ FTYKSSCAFS VFGTVCKFAC LAPFLLWLRK  SEG  ined sequences c  31   GAGGACACCC TTTGAGCAGC GGCTGCTG AATGCCTGCA AGAACAACA CATCACTCG GGCTGCTGCTAATC GCATCAAGC GTCAATCC GTCAATC GTAAGGAGC CCTGTAATC GTAAGGAGC GTTGAACTAC GTCATAAGAG GTTGAACTAC GTCATAAGAG GTTGAACTAC GTCATAAGAG GTCATAAAGG GTCACAAAAGG	OQRYTHLVAI EPNNRQKDED YTCKCDPGFS TMQCMSSGEW AQSLQCTSSG EQPAQVECT GFVLKGSKRL CEEGFELYGS PEGWTLNGSA CLRKAKKFVP  ID NO:237 PM2 OMESPOND to start  41  CAATGAGCCA TGATGGTGAA AAAGCCTTCTAAGGG GGGAACAACT TGACGGTGT TCAAGGGACT TCAAGGCACT AAGGAATTGCT TTTCTTCCC AAGAATTGCT TTTCTTCCCAAGAACAAACAAACAAACAAACAAACAAACA	ONKEETEYLM CVETYIKREK GLKCEQTVNC SAPIPACNVV NMDNEKPTCK TQGQWTQQIP QCGPTGEMIN TQLECTSQGQ ARTCGATCHW ASSCQSLESD  B DNA SEQUENCE and slop codons)  51 AAGGGGGTCC TATGCTAGAT ACTTGCCCAG GCTGATCCA TCTAGAAAAG ACTGGACCAT TCTAGAAAAG ACTGGACCAT TCTTTTAGAA TGCCTTGCGT TGCTTTTTTAGAA CAATGGTTCT TGAAAAGCAA AGAGTCAGAA CAATGGTTCT TGAAAAGCAA AGTGGACACA AGTGGACACA AGTGAACACC AGAAAGAATT	120 180 240 360 420 480 540 600 120 180 240 360 480 540 660 720 780 840 900

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                                                                             3000
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                                                                              600
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                                                                              720
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                                                                              780
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                                                                              900
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                                                                              960
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                                                                             1080
       LKRLNYDRKE LERRREASOH EIKDVLVWSN DRIIRWIQAI GLREYANNIL ESGVHGSLIA
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                                                                             1200
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       Nucleic Acid Accession #:
                                NM 016570
                                1-1134 (underlined sequences correspond to start and stop codons)
       Coding sequence:
80
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				GCGGATGTAT CCAACAGTAT			300 360
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				AGTACATCAA AGAATTCATG			480 540
	GTAGCAGGGA	ATTTTCACAT	AACAGTGGGC	AAGGCAATTC	CACATCCTCG	TGGTCATGCA	600
15				TACAATTTTT AATCCTTTAG			660 720
10				ATTACAGTTG			780
				TCTGTGACAG ATATTTATGA			840 900
20				TTCTGGCAGT			960
20				ATGTTACATG TCCTATAAAC			1020 1080
				CTTTTAGAAA			1000
25		PCI4 Protein segui					
	Protein Accession	1 #:	NP_057654				
	1	11	21	31	41	51	
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				ADVLDLAETM			120
			_	STSTALPPRE YNFSHRIDHL	-		180 240
35				SVTERERIIN MLHGIGKFIV			300 · · · · · · · · · · · · · · · · · ·
55	EDGHTDNHLP		1400112110	Mind Toke IA	BIICCKIKLO	SIVLANDALL	360
				SEC	ID NO:241 PBA7	DNA SEQUENCE	•
40	Nucleic Acid Acce	ession#: AA2191	134				
40					Coding se	quence: 2	24-1815 (underlined sequences correspond to start and stop codons)
	A ATTCCCCC	Г Т <i>С</i> СТТА АТТ	A ACCATOTTS	** ************************************	ro a trottotoa	CT GCTGCTGT	
4						CIT CAGATCA	
45						CTC GTCATTG AGA AGGACA	
						TC AGTTTATC	
						CTC TCTTCCAT	
50						AT TACGCATT	
						TG GGAGTTTT TG AAAGGAC	
				GT TAAGAGC			
		CATACITE A A TO	COTCAAACA	TO A ATATOAC			GAAC 000
55						TGG GATCTGT	TTC 720
55		A CAACATGCC	G ACCCGAAT	AA TGATAGG	ACT AACACT.		TTC 720 STAC 780
55	AAATCACTG( TTCAAAGCA	A CAACATGCC G CCAACCAA/ A TGAGGCAGC	G ACCCGAAT C ATATTGTT CT AGCCTCGC	TAA TGATAGG CT ATGCATCA CT CCACTGG	ACT AACACT. AC TGTTTTG. GGT TGGAGTC	TGG GATCTGT AGTA TTTTTTC AAG TCAGTTG GTC AAGGTCA	TTC 720 GTAC 780 GAT 840 ATTA 900
	AAATCACTGG TTCAAAGCAA GCACCATCCG	A CAACATGCC G CCAACCAA A TGAGGCAGC C TGCCACTCT	G ACCCGAAT AC ATATTGTT CT AGCCTCGC T CTTGTAGAC	TAA TGATAGG CT ATGCATCA CT CCACTGGC CC ATGTCGGC	ACT AACACT. AC TGTTTTG. GGT TGGAGTC AG CAAAACA	IGG GATCTGT AGTA TTTTTT AAG TCAGTTG	TTC 720 STAC 780 GAT 840 ATTA 900 TTG 960
55 60	AAATCACTGO TTCAAAGCAA GCACCATCCO GCTCCTCTGT TGAACTTCAO	A CAACATGCC G CCAACCAAA A TGAGGCAGC C TGCCACTCT GATGGCAGC C CCATATCTG	GG ACCCGAAT AC ATATTGTT TO AGCCTCGC T CITGTAGAC T TCGTTGGTC C AGAAGCCA	TAA TGATAGG CT ATGCATCA CT CCACTGGC CC ATGTCGGC IA CCATGGGC CA ATTCTATC	ACT AACACT. AC TGTTTTG. GGT TGGAGTC AG CAAAACA AT CGTAAAT AA CCAGTCC	TGG GATCTGT AGTA TTTTTTC AAG TCAGTTG CGTC AAGGTCA TTC CTCTGCA CTC AACATCC TTG GATGAGT	TTC 720 STAC 780 GAT 840 ATTA 900 TTG 960 ACA 1020 CTG 1080
	AAATCACTGC TTCAAAGCAA GCACCATCCC GCTCCTCTGT TGAACTTCAC TGATTTATGC	A CAACATGCC G CCAACCAAA A TGAGGCAGC C TGCCACTCT C GATGGCAGC C CCATATCTG G ACCAGGAAA	GG ACCCGAAT AC ATATTGTT CT AGCCTCGC T CTTGTAGAC T TCGTTGGTC C AGAAGCCA AC CTGTCAAC	TAA TGATAGG CT ATGCATCA CT CCACTGGC CC ATGTCGGC GA CCATGGGC CA ATTCTATC CA ACAACAA	ACT AACACT. AC TGTTTTG. GGT TGGAGTC AG CAAAACA AT CGTAAAT AA CCAGTCC TAC TCTCAGA	TGG GATCTGT AGTA TTTTTTC AAG TCAGTTG AGTC AAGGTCA TTC CTCTGCA CTC AACATCC	TTC 720 STAC 780 GAT 840 ATTA 900 TTG 960 ACA 1020 CTG 1080 AAAG 1140
	AAATCACTGC TTCAAAGCAA GCACCATCCC GCTCCTCTGT TGACTTCAC TGATTTATGG GGATTTCTTC GGGAGACGA	A CAACATGCC G CCAACCAAA A TGAGGCAGC C TGCCACTCT GATGGCAGC G CCATATCTG G ACCAGGAAA C CCATAGCAG C CTCAGCATC C CTCAGCATC	G ACCGAAT AC ATATTGTT T AGCCTCGC T CTTGTAGAC T TCGTTGGTCA AC CTGTCAAC A GCTCACC A GAGCCA C CTGTCAAC A GCTCACT C TTGCTAAA	TAA TGATAGG CT ATGCATCA CT CCACTGGC CC ATGTCGGC CA ATTCTATC CA ACACAA CA TGCCCTG TG CTGGATTA	ACT AACACT. AC TGTTTTG. GGT TGGAGTC AG CAAAACA AT CGTAAAACA AA CCAGTCC AA CCAGTCC AAC ACAGTCC AACACACACACACACACACACACACACACACACACA	IGG GATCTGT AGTA TTATTTT AGTA TCAGTTG AGTC AAGGTCA TTC CTCTGCA CTC AACATCC ITG GATGAGT AGAC CACTTCA GGG GATAAGA GGA TACCAG	TTC 720 GTAC 780 GAT 840 AATTA 900 TTG 960 ACA 1020 CTG 1080 AAAG 1140 AGAG 1200 ATAG 1260
	AAATCACTGC TTCAAAGCAA GCACCATCCC GCTCCTCTGT TGAACTTCAC TGATTTATGC GGATTTCTTC GGGAGACGA TCACAGACCC	A CAACATGCC G CCAACCAAA A TGAGGCAGC C TGCCACTCT C GATGGCAGC G CCATATCTG G ACCAGGAAA C CCATAGCAG C CTCAGCATC C TGGGGACG	G ACCGAATAC ATATTGTTT AGCCTCGCT T AGCCTCGCT T CTTGTAGAC T TCGTTGGTCC C AGAAGCCA C CTGTCAAC C AGCTCAAC C TTGCTAAC C CTGCCAAC C CTGCCAAC C CTGCCAAC C CTGCCAAC C CTGCCAAC C CCAGCTTT	TAA TGATAGG CT ATGCATCA CT CCACTGGC CC ATGTCGGC IA CCATGGGC CA ATTCTATC CA ACAACAA TACCCCTG TG CTGGATTA TT TGAAATGG TT TGAAATGG	ACT AACACT. AC TGTTTTG. GGT TGGAGTC AG CAAAACA AT CGTAAATCA CAAGTCC TAC TCTCAGA AG AAATGAT AAG CCACACT CT GTCCTTAA	IGG GATCTGT AGTA TTTTTTC AGG TCAGGTG COTC AAGGTC TTC CTCTGCA TTC AACATCC TTG GATGAGT IGGC CACTTC GTG GATAAGA GTG GATAAGA	TTC 720  GTAC 780  GAT 840  AATTA 900  TTG 960  ACA 1020  CTG 1080  AAAG 1140  AGAG 1200  ATAG 1250  TTG 1320
60	AAATCACTGC TTCAAAGCAC GCACCATCCG GCTCCTCTGT TGAACTTCAC TGATTTATGG GGATTTCTTC GGGAGACGA TCACAGACC TTTATGTTGG TCTTTCCTGG	A CAACATGCC G CCAACCAAA A TGAGGCAGC C TGCCACTCT GATGGCAGC C CCATATCTG G ACCAGGAAA C CCTCAGCATC C TGGGGACT T TGCTTTTTCA	GG ACCGAATAC ATATTGTTACAC ATATTGTAGAC T CITGTAGAC T TCGTTGGTC C AGAAGCCAA A AGCTCACT C TTGCTACAC A AGCTCACT C TTGCTACAC A AGCTCACT C TTGCTACA A AGCTCACT ATTGGTCTA A GGACGAGC A AGGACGAGC A AGGACGAGC	TAA TGATAGG CT ATGCATCA CT CCACTGGC CC ATGTCGGC A CCATGGGC CA ATTCTATC CA ACAACAA CA TGCCCTG TG CTGGATTA TGAAATGG G GACCAATGC CA TGCCTTG CA GCCATTG CA TGCATTA CA TGAATGC CA TGCCTTTA	ACT AACACT. ACT TGTTTTG. AGT TGGAGTC AG CAAAACA AT CGTAAAT AA CCAGTCC TAC TCTCAGA AG AAATGAT AG CACACT AGT TCCTTAGA AG CCACACT AGT TCCTTAGA AG CCACACT AGT TCCTTAGA AG CTTCTTAGC ACT TTCTTAGC	IGG GATCTGT AGTA TTTTTTT AGTA TCAGTTG AGTC AAGGTC TTC CTCTGCA TTC CTCTGCA TTG GATGAGT IGAC CACTTC GTG GATAAGA GAA TACCAG GAC AGCTTGC TTG CTCAGCGA ATG AACTGGG	TTC 720 STAC 780 GAT 840 AATTA 900 TTG 960 ACA 1020 CTG 1080 AAAG 1140 AGAG 1200 ATTA 1260 TTG 1320 AGA 1380 GGCA 1440
60	AAATCACTGG TTCAAAGCAA GCACCATCCC GCTCCTCTGT TGAACTTCAC TGATTTATGG GGATTTCTTC GGGAGACGA TCACAGACCC TTTATGTTGC TCAATCTCCTGG TCAATCTCCT	A CAACATGCC G CCAACCAAA A TGAGGCAGC C TGCCACTCT GATGGCAGC C CATATCTG G ACCAGGAAA C CCATAGCAG C CTCAGCATC C TGGGGACT T TGCTTTTTCA TGGGATCAG C CATCTCGCTT	G ACCGAAT AC ATATTGTT TT AGCCTCGC T CTTGTAGAC T TCGTTGGTC C AGAAGCCAA A AGCTCACT C TTGCTAAC A AGCTCACT C TTGCTAAA A CCAGCTTT ATTGGTCTAA AGCAGCGAGC A AGACGAGC A AGACGAGC A AGACGAGC A ACATTTTG	TAA TGATAGG CT ATGCATCA CT CCACTGGC CX ATGTCGGC CA ATCTATC CA ACAACAA CA TGCCCCTG TG CTGGATTA CT TGAAATGG G CA CCATGG CA TGCCCTG TG CTGGATTA CT TGAAATGG CA TGGCTTTA A CTGTAACTC	ACT AACACT. AC TGTTTTG. AG CAAAACA AT CGTAAAT AA CCAGTCC TAC TCTCAGA AG AAATGAT AAG CCACACT CC CTGCCTGA AG TCTTAGC AC TCTTAGC AC TCTTAGC AT TCTTATTG	IGG GATCTGT AGTA TITTITT AGTA TATTITT AGTA TAGTTG AGTC AAGGTC TTC CICTGCA TTC CICTGCA TTG GATGAGT IGAC CACTTC GTG GATAAGA TACCAG GCA GCTTGC TTG CTCAGCGA TTG CTCAGCGA TTG CTCAGCGA TTG CTCAGCGA	TTC 720 GTAC 780 GAT 840 AATTA 900 TTG 960 ACA 1020 CTG 1080 AAAG 1140 AGAG 1220 ATTAG 1260 TTG 1320 AGA 1380 GGCA 1440 GG 1500
60	AAATCACTGC TTCAAAGCAAC GCACCATCCC GCTCCTCTGT TGAACTTCAC TGATTTATGC GGATTTCTTC GGGAGACGA TCACAGACCC TTTATGTTGC TCATTCCTGG TCAATCTCCT TGTGCTTTAT TATATACAAT	A CAACATGCC G CCAACCAAA A TGAGGCAGC C TGCCACTCT GATGGCAGC C CCATATCTG G ACCAGGAAA C CTCAGCATC C TGGGGACGT TGCTTTTTCA TGGGATCAG C CATCTCGCT C ATTACAATT C CATGAGTCT TCATGAGTCT TCATGAGT TCAT	G ACCGAAT AC ATATTGTT TAGCCTCGC T CTTGTAGAC T TCGTTGGTC C AGAAGCCA A CCTGTCAAC C TTGCTAAC C TTGCTAAC C TTGCTAAC C TTGCTAAC C TTGCTAAC C CAGCTTT ATTGGTCTAC A GGACGAGC C ATGTTGC C ATGATTC C ATGGTCTA A GCATCCTC C ATGATCCTA A GCATCCTC	TAA TGATAGG CT ATGCATCA CT CCACTGGC CC ATGTCGGC CA ATTCTATC CA ACAACAA CA TGCCCCT TG CTGGATTA TT TGAAATGG G GACCAATGC CA TGGCTTTA A CTGTAACTC CA TTGTAACTC CA TGGATTA CTGTAACTC CA TTGTAACTC CA TTGTAACTC CA TTGTAACTC CA TTTTTGTTC CC TTTTTGTTC	ACT AACACT. AC TGTTTTG. GGT TGGAGTC AG CAAAACA AT CGTAAAT AA CCAGTCC TAC TCTCAGA AG CAACTCACTCAC AG TCTCAGA AG CCACACT ACT GTCCTTAC CC CTGGCTGG AC TTCTAGG GG CCTGCCAT GT TATGTTTA	IGG GATCTGT AGTA TITTITT AGTA TATTITTT AGTA TATTITTT TO TOTO TAGG TO TOTO TAGG TO TAGG	TTC 720  STAC 780  GAT 840  ATTA 900  TTG 960  ACA 1020  CTG 1080  AAAG 1140  AGAG 1200  ATTA 1260  TTG 1320  AGA 1380  GGCA 1440  GG 1500  TTA 1560  AAA 1620
60	AAATCACTGG TTCAAAGCAA GCACCATCCG GCTCCTCTGT TGAACTTCAC TGATTTATGG GGATTTCTTC GGGAGACGC TTTATGTTGC TCTTTCCTGG TCAATCTCCT TGTGCTTTAT AAGGGATGCTCT	A CAACATGCC G CCAACCAAA A TGAGGCAGC C TGCCACTCT GATGGCAGC C CCATATCTG G ACCAGGAAA C CCATAGCAGC C TCAGCATC C TGGGGACGT T TGGGGACGT TGGGTACAGC C TGAGCATC C TGCTTTTCA T TGGGATCAGC C TTTTGGAACACC C TTTTGGAACACC C TTTTGGAACAC	GG ACCGAATAC ATATTGTTACAC ATATTGTAGAC T COTTGTAGAC T TCGTTGGTCC C AGAAGCCAA A AGCTCACT CC TTGCTAAC A AGCTCACT CC AGAGCTAA CC CCAGCTTA A TCGGTCTAA A GGACGAGC G ACATTTTG C ATGGGTCTA A GCATCCCTA A GCATCCCTA A GCATCCCTA A AGATCCAT A ATATCAAT	TAA TGATAGG CT ATGCATCA CT CCACTGGC CC ATGTCGGC CA ATTCTATC CA TGCCCCTG TG CTGGATTA TGAAATGG G GACCAATGCCAATGCCAATGCCTT G CTGGATTA TGAAATGG CA TGGCTTTA A CTGTAACTC G ATCTTATT GG ATCTTATT GG AGCTAGCA	ACT AACACT. ACT TGTTTTG AGT TGGAGTC AG CAAAACA AT CGTAAAT AA CCAGTCC TAC TCTCAGA AG AAATGAT AAG CCACACT CC TGGCTGG AC TTCTAGC AG TCTTATTG AG TCTTATTG AG TCTTATTG AG TCTTTATTG AAA AGTGAA	IGG GATCTGT AGTA TTATTTG AAG TCAGTTG CATC AAGGTCA CTC AACATCCA CTC GATGAGT AGAC CACTTCA GTG GATAAGA GTA TACCAGA GCC AGCTTGC TTG CTCAGCGA ACTGGC ACCCGGA ACTGGC GG GTGTGCCTTGC GG GTGTGCCTTGC GG GTGTGCCTTGC GG GTGTGCCTTGC GG GTGTGCCTTGCCT	TTC 720  STAC 780  GAT 840  AATTA 900  TTG 960  ACA 1020  CTG 1080  AAAG 1140  AGAG 1200  ATTA 1260  TTG 1320  AGA 1380  GGCA 1440  GG 1500  TTA 1560  AAACA 1620  AACA 1680
60	AAATCACTGG TTCAAAGCAA GCACCATTCAC GCACCATCCC GCTCCTCTGT TGAACTTCAC TGATTTATGG GGATTTCTTC GGGAGACGA TCACAGACCC TTTATGTTGC TCATTTCCTC TGTGCTTTAT TATATACAAT AGGGATGCTT AACCCCAGG	A CAACATGCC G CCAACCAA A TGAGGCAGC C TGCCACTCT GATGGCAGC C CCATATCTG G ACCAGGAAA C CTCAGCATC C TGGGGACGT TGGGTTTTTCA TGGGATCAG C CATCAGCATC C TGGGATCAG T CATCAGCATC C TTTTGGAACA T TGGATCAG T TGGATCAG T TATGAGTCT ATATACATC TATGAGTCT ATATACAGTCT ATATACAGTCT ATATACAGTCCT ATATACAGTCCT ATATACAGTCCT AGCAGCTCTT AGGGCTCT AGGCTCTT	G ACCGAAT AC ATATTGTT T AGCCTCGC T CTTGTAGAC T TCGTTGGTC C AGAAGCCAC C TTGCTAAA C CTGTCAAA C CCAGCTTT A TTGGTCTAAA C CAGCTTT A TTGGTCTAA A GGACGAGC G ACATTTTG C ATGAGTCTA A GGACGAGC T ATATCAAT T CACCAAGAA T CACCAAGAA T CACCAAGAA T G GAGTGTAA	TAA TGATAGG CT ATGCATCA CT CCACTGGC CT ATGCTGGC CA ATTCTATC CA ACAACAA CT TGAAATGG CT CTGATT CT TGAAATGG CT TTTTATT CT TGAAATGG CT TTTTTTTT CT TGAATTATT CT TGAATTATT CT TGAATTATT CT TGAATTATT CT TTTTTTTT CT TTTTTTTT CT TTTTTTTT CT TAGTT CA AGCTTGT CA AGCTTGT CT ATGCTTGTT CA AGCTTGT CA AGCTTGT CT ATGCTTGTT CA AGCTTGT CA AGCTTGT CA AGCTTGT CT TTTTTGTT CA AGCTTGT CA AGCTTGT CT TGTT CT TGT CT	ACT AACACT. AC TGTTTTG AG CAAAACA AT CGTAAAT AA CCAGTCC TAC TCTCAGA AG AAATGAT AAG CCACACT AC TCTCAGA AC TCTCAGA AC TCTTAGC AC TTCTAGC AC TTCTAGC AC TTCTAGC AC TTCTAGC AC TTTATTGG AC CTGCCAT AC CTGCCAT AC CTGCCAT AC CTGCCAT AC TCTAGC AC CTGCCAT AC CC AAAACAG CC AAAACAG CG TAGGGG	IGG GATCTGT AGTA TTTTTTT AGTA TCAGTTG AGTC AAGGTCA CTC AACATCC ITG GATGAGT GGAC CACTTCA GGAC CACTTCA GGAC AGCTTGC ATG GATAGAG AGAA TACCAG AGC AGCTTGC ATG GACTGG ACTGGAC ACTGGAC ACTGCATG ACTGGAC ACTGCATG ACTGACAC ACTGACAC ACTGACAC ACTCAAAAAA CCCAAAAAAA	TTC 720 GTAC 780 GAT 840 AATTA 900 TTG 960 ACA 1020 CTG 1080 AAAG 1140 AGAG 1200 ATTAG 1260 TTG 1320 AGAG 1380 GGCA 1440 GG 1500 FTA 1560 AAAC 1680 AACAA 1680 AACAA 1740 CCAGC 1800
60 65 70	AAATCACTGG TTCAAAGCAA GCACCATCCC GCTCCTCTGT TGAACTTCAC TGATTTATGG GGATTTCTTG GGAGACGA TCACAGACCC TTTATGTTGC TCATCCCT TGTGCTTTAT TATATACAAT AGGGATGCTT ACATTTGTTT AACCCCAGG TTTCTCCAGA	A CAACATGCC G CCAACCAAA A TGAGGCAGC C TGCCACTCT GATGGCAGC C CCATATCTG G ACCAGGAAA C CTCAGCATC C TGGGGACGT C TGGGGACGT C TGTTTTCA T TGGGATCAC C TATACAATT C CATGAGTCT C TTTTGGAACA T TTTGGAACA T TTTGAGTCAT A GCAGCTCT A GCAGCTCT A GCAGCTCT A GCAGCTCT A GCAGCTCT A GCACCTAATG	G ACCGAAT AC ATATTGTT TT AGCCTCGC T CTTGTAGAC T TCGTTGGTC C AGAAGCCA ACCTGTCAAC C TTGCTAAA C CAGCTTA A TGGTTCAAC A AGCTCACT C TTGCTAAA A GACGCACT ATTGGTCTA A GGACGAGC A AGTTTTG C ATGAGTCTA A GCATCCCT A ATATCAAT T CACCAAGA G GAGTGTAA G GAGTGTAAA G CCTCAACA	TAA TGATAGG CT ATGCATCA CT CCACTGGC CC ATGTCGGC CA ATTCTATC CA ACAACAA TG CTGGATT TG CTGGATT TG TGGATT TGAAATGG GACCAATG CA TGCTTATC CA TGCTTATC CA TGCTTATC CA TGCTTATC CA TGCTTATC CA TGCTTATC CC TTTTTGTT CG ATCTACT CA AGCTGT CC TTCTGAAC CC TTCTGAAC CC TGCTTAAC CC AGCTGT CC TTCTGAAC CC TTCTGAAC CC TTCTGAAC CC TTCTGAAC	ACT AACACT. AC TGTTTTG. GG TGAAATCAAAT CGTAAATCAACT. AAC CCAGTCC TAC TCTCAGA AC CCACACT. AC TCTCAGA AC CTGCCTTAC AC TCTCAGA AC TTCTAGA AC CTGCCTGG GG CCTGCCAT TATGTTA AAA AGTGAAA ACC AAAACAC ACC AAAACAC ACC AAAACAC ACC AAAACAC ACC GG TGGCTGG ACT TGTGTTA	IGG GATCTGT AGTA TTTTTTT AGTA TTATTTT AGTA TCATTGT AGTC AAGGTCA TTC CTCTGCA TTC CTCTGCA TTC GATGAGT IGAC CACTTCA GTG GATAAGA TGAA TACCAG GCC AGCTTGC TTG CTCAGCGA ATG AACTGGG GC CTGCCATG GG GTGTGCTT ITA CCTGAGAA CCTCAAAAAA	TTC 720  STAC 780  STAC 780  GAT 840  ATTA 900  TTG 960  ACA 1020  CTG 1080  AAAG 1140  AGAG 1200  ATTA 1320  AGA 1380  GGCA 1440  GG 1500  TTA 1560  AAAC 1620  AAACA 1680  AACA 1680  ACAGA 1740  ACAGC 1800  CTTA 1860
60	AAATCACTGG TTCAAAGCAA GCACCATTCC GCTCCTCTGT TGAACTTCAC GGATTTCTTC GGGAGACGA TCACAGACCC TTTATGTTGC TCATTCCTGG TCAATCTCCT TGTGCTTTAT TATATACAA AGGAGATCT ACATTGTTT AACCCCAGG TTTTCTCCAGG GAACTGGTTT	A CAACATGCC G CCAACCAA A TGAGGCAGC C TGCCACTCT GATGGCAGC C CCATATCTG G ACCAGGAA C CTCAGCATC C TGGGGACGT C TGGGGACGT C TGGGGACGT C TGGGGACGT C TGTGGACA T TGGGATCAG C TTTTGGACA T CATGAGTCT C TTTGGAACA T TATGAGTCA C TGGACCTAATG C TTTGGAACA T TGAAGAGAC C TTTGAACA T TGAAGAGAC C TTTGAACAA T TGAAGAGAC	G ACCGAAT AC ATATTGTT TT AGCCTCGC T CTTGTAGAC T TCGTTGGTC C AGAAGCCAA A AGCTCACT C TTGCTAAC A AGCTCACT ATTGGTCTAA A GCACGAGC G ACATTTTG C ATGAGTCTA A GCATCCCT C ATGAGTCTA A GCATCCT C ATGAGTCTA C CCCAAGAA G CCTCAACAC C AGCTCAACAC C AGCTCAACAC C AGCTCAACAC C ACCTCAACAC C A CTCCAACAC C A CTCCAACAC C A CTCCAACAC C A CTCCATCAACAC C A CCCTCAACAC C A CTCCTGAAA	TAA TGATAGG CT ATGCATCA CT CCACTGGC CX ATGTCGGC CA ATCTATC CA TGCCCTG TG CTGGATTA TT TGAAATGG G GACCAATG G GACCAATG G ATCTTATC GG ATCTTATC GG AGCTAGC CA GACTAGC CA GCACTCCT TG CGACTCCT TG ATAAAGA	ACT AACACT. ACT TGTTTTG. AGT TGGAGT. AG CAAAACA AT CGTAAAT. AA CCAGTCC. TAC TCTCAG. AG AAATGAT. AG CCACACT. AG CCTGGCTGG. AC TTCTAGC. AA TCTTATTG. AAA AGTGAA. ACT AGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGG	IGG GATCTGT AGTA TTTTTTT AGTA TTATTTT AGTA TTATTTT AGTA TAGTA AGT CAGTCA TTC CTCTGCA TTC CTCTGCA TTC AACATCC TTG GATGAGT AGAC TACATT AGAC CACTTCA GTG GATAAGA TGAA TACCAG GCC AGCTTGC TTG CTCAGCGA ATG AACTGGG GC CTGCCATG GG GTGTGCTT IA CCTGAGAC CTAT GTGAAA CCAA TACAAG CCT CAAAAAA CCAA TCCAGG TTT CAGTGTCA TTT CAGTGTCA TTT CAGTGTCA TTCC CCCTCCT	TTC 720 GTAC 780 GAT 840 AATTA 900 TTG 960 ACA 1020 CTG 1080 AAAG 1140 AGAG 1200 ATTAG 1260 TTG 1320 AGA 1380 AGCA 1440 AGG 1500 CTA 1560 AAAAG 1680 AAAAG 1740 CCAGC 1800 CTTA 1860 ATTAG 1860 ATTAG 1860 ATTAG 1920 CCMC 1980
60 65 70	AAATCACTGG TTCAAAGCAA GCACCATTCCG GCACCATCCC GCTCCTCTGT TGAACTTCAC TGATTTATGG GGATTTCTTGG GGAGACGA TCACAGACCC TTTATGTTGC TCATTTCCTGG TCAATCTCCT TGTGCTTTAT TATATACAAT AGGGATGCTT AACCCCAGG TTTCTCCAGA GGAGGGTGTT CAGAAGGAA	A CAACATGCC G CCAACCAA A TGAGGCAGC C TGCCACTCT GATGGCAGC C CCATATCTG G ACCAGGAAA C CTCAGCATC C TGGGGACGT TGGGTTTTTCA TGGGATCAG T CATGAGTCT ATATACAATT C ATGAGTCT ATATACAATT A GACCTCT A GACCTAATG C TTTGGACCA C TTTGGACCA C TTTGGACCA C TTTGGACCA C TTTGGACCA C TTTGGACCA C TTTGACCAC C TTTAGACCAC C TTAAAAAGCA C C CTCAAAAAGCA	G ACCGAAT AC ATATTGTT TAGCCTCGC T CTTGTAGAC T TCGTTGGTC C AGAAGCCA C CTGTCAAC C TTGCTAAA C CCAGCTTT A TGGTCAA A GGACGAGC G ACATTTTG C ATGAGTCT A ATATCAAT T CACCAAGAA G GAGTCAAC A GGAGTCAAC A TGATGATAA T CACCAAGAA T CACCAGAAA T CACCAAGAA T CACCAAGAAA T CACCAAGAA T CACCAA	TAA TGATAGG CT ATGCATCA CT CCACTGGC CT ATGCTGGC CA ATTCTATC CA ACAACAA CT CT CCACTGGC CA ATTCTATC CA ACAACAA CT C	ACT AACACT. ACT TGTTTTG AG CAAAACA AT CGTAAAT AA CCAGTCC TAC TCTCAGA AG AAATGAT AAG CCACACT ACT GTCCTTAGC ACTTTTTAGC ACTTTTTAGC ACTTTATTGG GCCTGCCAT AAAAAAAAAA	IGG GATCTGT AGTA TTTTTTT AGTA TTTTTTT AGTA TTATTTT AGTA TCAGTTG AGTC AAGGTC ATC CTCTGCA TTC CTCTGCA TTG GATGAGT IGAC CACTTC AGTG GATAAGA TACCAGG AGC AGCTTGC TG CTCAGCGA ATG AACTGGG GC CTGCCATG TG GTGTGCT TA CCTGAGAC CTAT GTGAAA CCT CAAAAAA CCT CAAAAAA CCT CAAAAAA CTC CAAAAAA CTC CAAAAAA TCAGG GC CTGCC CTGC CTCT CAGGAC CTCT CAGAAAAA CTC CAAAAAA CTC CAAAAAA CTC CAAAAAA CTC CAAAAAA CTC CAGAACAC CTT CAGTGTC	TTC 720  STAC 780  STAC 780  GAT 840  ATTA 900  TTG 960  ACA 1020  CTG 1080  AAAG 1140  AGAG 1200  ATAG 1260  TTG 1320  AGA 1380  GGCA 1440  GG 1500  TTA 1560  AAA 1620  AAAA 1680  AGAA 1740  CCAGC 1800  CTTA 1860  ATG 1920  CCMC 1980  GAGC 2040
60 65 70	AAATCACTGG TTCAAAGCAA GCACCATCCC GCTCCTCTGT TGAACTTCAC TGATTTATGC GGATTICTCC GGGAGACGA TCACAGACCC TTTATGTTGC TCATTCCTCG TCATCTCCT TGTGCTTTAT TATATACAAT AGGGATGCTT AACCCCAGG TTTCTCCAGA GGAGGGTGTT CAGAAGGAA AGGATATCAC AGAAGGAA AGGATATCAC AGAGCAGCC	A CAACATGCC G CCAACCAAA A TGAGGCAGC C TGCCACTCT GATGGCAGC C CATATCTG G ACCAGGAAA C CATAGCAG C CTCAGCATC C TGGGGACGT T TGCGATCAGC C TTTGGATCAG C TTTGGAACA T TATACAATT C TGGGACCT C TTTGGAACA T TTGAACAC C TTGAACAGC C TTTGAACAG C TTTAAAAAG T TTGAACAAAAG T TTGAACAAAAAA T TTGAATAGA	GG ACCGAATAC ATATTGTTAGAC ATATTGTTAGACACT TCGTTGGACACACACACACACACACACACACACACACACA	TAA TGATAGG CT ATGCATCA CT CCACTGGC CX ATGTCGGC CA ATGTCTGGC CA ACATGGC CA ACAACA CA TGCCCTG TG CTGGATTA TT TGAAATGG CA TGGTTTA A CTGTAACT GG ATCTTATT GG ATCTTATT GG AGCTAGC AG CATTAGT CA AGCTGG CT TTTGTA CC TTCTGAAC CC TTCTGAAC CC TCTGAAC CCT TGGAAC CCT TGGAAGAC CCT TGAAGAC CCT TGAAGAC	ACT AACACT.  ACT TGTTTTG AGT TGGAGT AG CAAAACA AT CGTAAAT AAA CCAGTCC TAC TCTCAGA AG AAATGAT AAG CCACACT CC CTGGCTGG AC TTCTAGC AT TCTTATTG AG TCTTATTG AG TCTTATTG AG TCTTATTG AG TAGGGT AGT TAGGCAT	IGG GATCTGT AGTA TITTITT AGTA TITTITT AGTA TAGTTG AGTC AAGGTCA TTC CTCTGCA TTC CTCTGCA TTG GATGAGT IGAC CACTTCA GTG GATAAGA TGAA TACCAG GGC AGCTTGC TG CTCAGCGG TG CTCAGCGG TG GTGTGCT IA CCTGAGAC CTAT GTGAAA CCT CAAAAA CCA TCAAGG GCC AGAACAC TTT CAGTGTCC TTC CCTCCT TTCCTCCT CTCT TITTCTC	TTC 720  STAC 780  GAT 840  AATTA 900  TTG 960  ACA 1020  CTG 1080  AAAG 1140  AGAG 1200  ATAG 1260  TTG 1320  AGA 1380  GGCA 1440  GG 1500  FATA 1560  AAAC 1680  AGAA 1740  CCAGC 1800  CTTA 1860  ATG 1920  CCAC 2100  FATG 2160

ACACACAGTG TGGACAACTG CCCATATATT CTATCTAGAT TAGGAGAGGG TCCTGGCTAG 2340 GATTTTAGTG GTAATTCCTA GTTACATTCA ACAAGTATAA AGATTATAGA GCTTATTTTA 2400 TGAACTATAA ACTATAATTT AATGCAAAAT ATCCTTTTAT GAATTTCATG TTAATATTGT 2460 GAAATATTAA AATAATTCCR CAATAGTTGA GAAAAATGAG CATTTTTTC CATTTTTAAA 2520 5 AAATGCATAG AAAAGACAAT TTTAAAATCC TGGGACCATA TTTATTTAGA AGTAGCTGTT 2580 AGTAAAACAT TAGAAAAGGA GTCAGGCCAT TAGGTTATTT ATCCAAATCT CTAAGCAATT 2640 AGGTTGAAGT TATTAAGTCA AGCCTAGAAA AGCTGCCTCC TTGTAAGGCT TTCATGACAA 2700 TGTATAGTAA TCCACAGTGT CCAATTCTTC ACACTCCTCA GGAATATCAC TACCTCAGGT 2760 TACGGTACAC AGGCTATAAT TGATGATGAT GTTCAGATAA CTGAAGACAC AATAAATGAC 2820 10 ATTCAGACAT CAGGAMAAWW CCCTCATGTT CTTTTCTATG ATGGCCACCT GTACCAGCAA 2880 CGTGGGTTTC ACCCACACAA CGATGAACTG TTCTCTTACT TCTCCAGTTG ATTTTAAAGA 2940 CTTGTTAAGA GGTCTTACTA ATAAAATTTG GGTATGATAG AAAAWCCACA ATCAAAWCTT 3000 GAACCAAATA ACATATTAAA TTACTAATAT TTAAGTGATG GAAGACACAC AAAAAACTTA 3060 AAAGCACGAA CAACCTAACT TGAAAAAGAA TTTTAAAATA TGATTAACCT GAAGAAAAGA 3120 15 GAATCCTAAG AGCCAAAGCT CCTTTTTATT TAGCTTGGAA TTTTCCTATT GGTTCCTAAC 3180 AAACTGTCCC AATGTCATAT AAGGAAACAT GATCTATTAC ATTCCTTTAT AACAATGTGG 3240 AGAGACTATA AACCTATGTA AGTAGTAAAA CTATATYAGA GACTCAGGAG ACTGACTAAA 3300 AGGCCTGGAT CTGCAGTGTA TTATCTGTAT AAAAATTGGC AGGGGGAAGC TAAAAGGAAA 3360 GGAGATTGGA GATCTCAATT CTATCATGGT GTATTTCATA CGCAAATCAG AGCATGCATT 3420 20 GTTTTTGTT TTTGGAAAGA GAAGGGAAGT GTGTTCTGCC CCATGTTTCC TTCCGTGTTT 3480 ATAGTTCAAA CTCTATATAT ACTTCAGGTA TTTTTTGTTT AGCCCTTCAT TATAAATGGG 3540 CAGGAAATTG TTTATCAACC TAGCCAGTTT ATTACTAGTG ACCTTGACTT CAGTATCTTG 3600 AGCATTCTTT TATATTTTTC TTTTATTATC CTGAGTCTGT AACTAAACAA TTTTGTCTTC 3660 AAATTITTAT CCAATATCCA TIGCACCACA CCAAATCAAG CTTCTTGATT TICAAAAAATA 3720 AAAAGGGGGA AATACTTACA ACITGTACAT ATATATTCAC AGTTTTTATT TATAAAAAAA 3780 25 ATTTACAGTA CTTATGGAGA GCCAGCAGAA GACATCAGAG CACTCACTTC TTCCCATCTT 3840
TGTTAAGGTT AGCGAATTAC CCATGGACAC TGTTAGGTGA GGCTCATTCG GCAGCCCTGA 3900 AAACAAACCT GGTCACACTG TCTTTACCCT CTCCCTTCAG ATAAAGCACT TCGATTATCT 3960 ATTGATCTGC CCAGTTTTCA AGTCATGCGA ATACTAAAAA GGTTACATCA TCTGGATCTG 4020 30 TACCTTGGCT ATATAAGCAT GTTTTCCCCC TATTCTATGT TTCTTTTTTT GGTGAACATT 4080 GAAAAACAGG AGGTGACTTA TTACTGTTAA TTAAAACTAA ATGAAAAATG TCAAGTCTTT 4140 AAAACAGTGA GCTTGTAACT CTTTCATGTA ATTTTATTCT CTATGAATTT GGCTATCCTA 4200 CTGAATCTTA AAATAAAGGA AATAAACACT TTTTTTTWAA AAAAAAGGAA AAATAMAARW 4260 MWAAAAATCT CAATGAAATA TTTCACAAGA AGGAAAAA 35

SEQ ID NO:242 PBA7 Protein sequence:

#### AAF91431 Protein Accession #:

40 MFTFLSSVTA AVSGLLVGYE LGIISGALLQ IKTLLALSCH EQEMVVSSLV IGALLASLTG 60 GVLIDRYGRR TAIILSSCLL GLGSLVLILS LSYTVLIVGR IAIGVSISLS SIATCVYIAE 120 IAPQHRRGLL VSLNELMIVI GILSAYISNY AFANVFHGWK YMFGLVIPLG VLQAIAMYFL 180 PPSPRFLVMK GQEGAASKVL GRLRALSDTT EELTVIKSSL KDEYQYSFWD LFRSKDNMRT 240 RIMIGLTLVF FVQITGQPNI LFYASTVLKS VGFQSNEAAS LASTGVGVVK VISTIPATILL 300
VDHVGSKTFL CIGSSVMAAS LVTMGIVNLN IHMNFTHICR SHNSINQSLD ESVIYGPGNL 360
STNNNTLRDH FKGISSHSRS SLMPLRNDVD KRGETTSASL LNAGLSHTEY QIVTDPGDVP 420
AFLKWLSLAS LLVYVAAFSI GLGPMPWLVL SEIFPGGIRG RAMALTSSMN WGINLLISLT 480
FLTVTDLIGL PWVCFIYTIM SLDLIGLPWV CFIYTIMSLA SLLFVVMFIP ETKGCSLEQI 540 45 SMELAKVNYV KNNICFMSHH QEELVPKQPQ KRKPQEQLLE CNKLCGRGQS RQLSPET 50

#### SEQ ID NO:243 PAB4 DNA sequence: Nucleic Acid Accession#: AA172056

55

75

Coding sequence:

121-339 (underlined sequences correspond to start and stop codons)

TTTAGCCACC AGAGGANTTC TCTTGAAATA CCCAAAATCC ATCAGTATCT TGAATCATGC 60 TGGATTTTGA AGAATTCTTA AGAAGCCATG TAAAGGGGGC TCTCTGGCCT TGAAATAGTG 120 ATGTTTTTTA TACAGAAAGG AGAATGCAGA ATGGTCAGAC TATCATGCAC TGTTAAATTT 180 60 GATTICAAGA AATTACAGGA AAACTTICCA AAGTTCCATC TCACAGAANN TTATTTINCC 240 AAGAATTCCA AGATAAGTTT AGTTTTATGG AAGACTTTTA TGTGGTTTTT ACTCACTCTT 300 CATCTCAGAC ATCGACAGAT GATTACATCA CITATAGTTC TAGTAAATTT ATTAATATAA 360 AACTCAGAGA CATTCCAATA TCCACATTGC TTACACCATT AGGCATAGAT TCAGTGTCAG 420 CTATGACAAT TGAAAATGAG CTGTTTTGTG ATTTAAAGGT TTAAATTTCT CTAACCAAAC 480 65 TGCTTGATCC AGATGCAGGA CTGCAAATGT TAATATTTGT TCTGGAAGAA CAATCAAATA 540 AGACTIAAGA GGAAAGGAA TIGCCACAAT CCACCTGAAA TITITTCTTA AAAATCAATA 540
AGACTIAAGA GGAAAGGGAA TIGCCACAAT CCACCTGAAA TTTITTCTTA AAAAGTGTGC 600
AGCCTACTAA ATCAGAATGA AAATAGAAGT ACAAGATTAT AAAACAAAATG CAATCAAACT 660
TTTCTTAAGC TTACCTAAAG TTATTTCATC TIGAAAATTTC AAGCAACTTT GTTCAACATT 720
AAATTGACAA TCTAAACTAA CAAGTCTTTT GAATTTTATGC ATGGTAGTAA ACATTCTCTC 780
TATTAACTTT ATTACCTAAG GCTAAACCTA AAATTTTAA GCAAAATTAG AAAAATAGTC 840
TTCACTCATC AAAAATAAA GTTTGTTACA TTTAGTATTT TCCCAATAAAA ATTGGTCGTT 900 70 CTTGGTTTTT TATTTGGAGA GTCTGTGCAA AATGTCACTA AAAATAAATT AGCACTAGAA 960 ATTATTTCTA AATACCAAA

SEQ ID NO:244 PBQ8 DNA SEQUENCE

Nucleic Acid Accession#: X51405 Coding sequence:

3-1721 (underlined sequence corresponds to start and stop codon)

1 11 51 21 31

	1	ı	1	1	ı	1	
	AAATGGCGTG	CCCGTCTCTC	cccccccc	CTGCCTCGCA	GTGGTTTCTC	CTGCAGCTCC	60
			GTGCAGCCCG				120
			GACACTCATT				180
5			CAGCAGCGCC				240
_	AAGAGGCCGC	CCGCGTAGGA	AGGCACGGCC	GCCGCCGCC	GAGCGCAGCG	ATGGCCGGGC	300
	GAGGGGGCAG	CGCGCTGCTG	GCTCTGTGCG	GGGCACTGGC	TGCCTGCGGG	TGGCTCCTGG	360
			GGGGCGCCCG				420
	AGCAAGAGGA	CGGCATCTCC	TTCGAGTACC	ACCGCTACCC	CGAGCTGCGC	GAGGCGCTCG	480
10	TGTCCGTGTG	GCTGCAGTGC	ACCGCCATCA	<b>GCAGGATTTA</b>	CACGGTGGGG	CGCAGCTTCG	540
	AGGGCCGGGA	GCTCCTGGTC	ATCGAGCTGT	CCGACAACCC	TGGCGTCCAT	GAGCCTGGTG	600
	AGCCTGAATT	TAAATACATT	GGGAATATGC	ATGGGAATGA	GGCTGTTGGA	CGAGAACTGC	660
	TCATTTTCTT	GGCCCAGTAC	CTATGCAACG	AATACCAGAA	GGGGAACGAG	ACAATTGTCA	720
	ACCTGATCCA	CAGTACCCGC	ATTCACATCA	TGCCTTCCCT	GAACCCAGAT	GGCTTTGAGA	780
15	AGGCAGCGTC	TCAGCCTGGT	GAACTCAAGG	ACTGGTTTGT	GGGTCGAAGC	AATGCCCAGG	840
	GAATAGATCT	GAACCGGAAC	TTTCCAGACC	TGGATAGGAT	AGTGTACGTG	AATGAGAAAG	900
	AAGGTGGTCC	AAATAATCAT	CTGTTGAAAA	ATATGAAGAA	AATTGTGGAT	CAAAACACAA	960
	AGCTTGCTCC	TGAGACCAAG	GCTGTCATTC	ATTGGATTAT	GGATATTCCT	TTTGTGCTTT	1020
	CTGCCAATCT	CCATGGAGGA	GACCTTGTGG	CCAATTATCC	ATATGATGAG	ACGCGGAGTG	1080
20			TCCTCCCCAG				1140
	CATACTCTTC	TTTCAACCCG	GCCATGTCTG	ACCCCAATCG	GCCACCATGT	CGCAAGAATG	1200
			GATGGAACCA				1260
			TACCTTAGCA				1320
05			GAAGAGACTC				1380
25			CAGATACACC				1440
			GCCACCATCT				1500
			TGGAGATTGC				1560
•			ATAACAAAGA				1620
20			TCATTTTCTG				1680
30			TCAGAAACTT				1740
			TGTAGTATGA				1800
			TTTATTTTTT				1860
			AAAAATATAA				1920
35			TTACACAAAA				1980
33			ATTCCTGGTA				2040
			GAAGTTCTTT				2100
			CAGATACAGC				2160 2220
			GAAGAAAAGG				2220
40			TTGTACATAT				2340
70			AGGGTTTTCT				2400
			AAAAAATCCC			CANTIOCALL	2400
	CIGNATONAL	MAN I DUNN	and the same of th	C-31GRANAN	en mi	*	

## SEQ ID NO:245 PBQ8 Protein sequence P16870 45

Protein Accession#:

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MAGRGGSALL ALCGALAACG WILLGAEAQEP GAPAAGMRRR RRLQQEDGIS FEYHRYPELR 60
EALVSVWLQC TAISRIYTVG RSFEGREILLV IELSDNPGVH EPGEPEFKYI GNMHGNEAVG 120
RELLIFLAQY LCNEYQKGNE TIVNLIHSTR IHIMPSLNPD GFEKAASQPG ELKDWFVGRS 180
NAQGDLNRN FPDLDRIVYV NEKEGGPNNH LLKNMKKIVD QNTKLAPETK AVHIWIMDIP 240
FVLSANLHGG DLVANYPYDE TRSGSAHEYS SSPDDAIFQS LARAYSSFNP AMSDPNRPPC 300
RKNDDDSSFV DGTTNGGAWY SVPGGMQDFN YLSSNCFEIT VELSCEKFFP EETLKTYWED 360
NKNSLISYLE QHRGVKGFV RDLQGNPIAN ATISVEGIDH DVTSAKDGDY WRLLIPGNYK 420
LTASAPGYLA ITKKVAVPYS PAAGVDFELE SFSERKEEEK EELMEWWKMM SETLNF

SEQ ID NO:246 PBY4 DNA sequence Nucleic Acid Accession#: AF038966

	HUCKEL AND ALCOSINIA.		W. 600300					
		Codin	g sequence:	91-1107 (unde	riined sequence c	orresponds to start	and stop codon)	
60								
	1	11	21	31	41	51	•	
	1	1			1	1		
	GGGGCGACGT	GAGCGCGCAG	GGGGGCGCG	GCCTCGCCTC	GTCTCTCTCT	CTGCGCCTGG	60	
	GTCGGGTGGG	TGACGCCGAG	AGCCAGAGAG	ATGTCGGATT	TCGACAGTAA	CCCGTTTGCC	120	
65	GACCCGGATC	TCAACAATC	CTTCAAGGAT	CCATCAGTTA	CACAAGTGAC	AAGAAATGTT	180	
	CCACCAGGAC	TTGATGAAT	TAATCCATTC	TCGGATTCTA	GAACACCTCC	ACCAGGCGGT	240	
	GTGAAGATGC	CTAATGTAC	CAATACACAA	CCAGCAATAA	TGAAACCAAC	AGAGGAACAT	300	
	CCAGCTTATA	CACAGATTG	AAAGGAACAT	GCATTGGCCC	AAGCTGAACT	TCTTAAGCGC	360	
	CAAGAAGAAC	TAGAAAGAA	AGCCGCAGAA	TTAGATCGTC	GGGAACGAGA	AATGCAAAAC	420	
70	CTCAGTCAAC	ATGGTAGAA	AAATATTTGG	CCACCTCTTC	CTAGCAATTT	TCCTGTCGGA	480	
	CCTTGTTTCT	ATCAGGAAT	TTCTGTAGAC	ATTCCTGTAG	AATTCCAAAA	GACAGTAAAG	540	
	CTTATGTACT	ACTIGIGGA:	GTTCCATGCA	GTAACACTGT	TTCTAAATAT	CTTCGGATGC	600	
	TTGGCTTGGT	TTTGTGTTG	* TTCTGCAAGA	GCGGTTGATT	TTGGATTGAG	TATCCTGTGG	660	
	TTCTTGCTTT	TTACTCCTT	TTCATTTGTC	TGTTGGTACA	GACCACTTTA	TGGAGCTTTC	720	
75	AGGAGTGACA	GTTCATTTAC	ATTCTTTGTA	TTCTTCTTCG	TCTATATTTG	TCAGTTTGCT	780	
	GTACATGTAC	TCCAAGCTG	: AGGATTTCAT	AACTGGGGCA	ATTGTGGTTG	GATTTCATCC	840	
	CTTACTGGTC	TCAACCAAA	TATTCCTGTT	GGAATCATGA	TGATAATCAT	AGCAGCACTT	900	
	TTCACAGCAT	CAGCAGTCAT	CTCACTAGTT	ATGTTCAAAA	AAGTACATGG	ACTATATCGC	960	
00			GAAGGCCCAA				1020	
80	AAAACTGTCC	AGACCGCAG	TGCAAATGCA	GCTTCAACTG	CAGCATCTAG	TGCAGCTCAG	1080	

	AATGCTTTCA	AGGGTAACCA	GATTTAAGAA	TCTTCAAACA	ATACACTGTT	ACCTTTTGAC	1140
	TGTACCTTTT	TCTCCAGTTA	CTGTATTCTA	CAAATATTTT	TATGTTCAAA	ACACACAGTA	1200
<b>,</b>	CAGACAGCAT	GGATATTTCC	TGTTCACTTG	TGCATGGGCT	AAAACCAGGA	AAACTTCCTT	1260
	GTCTTATTAC	TTTACCTAAT	AGTTTCTTAA	TATTTCAGTG	CCCCTTGCAG	TTATAAAAAA	1320
5	ACATGCTAAA	TAAATATTCT	CCATATTTTT	GGGGGATGAC	ATTCAGTGAA	TTATTTCAGT	1380
	GGTGACCCAC	TGAAAATTAA	TAATGGTACT	TATGATTAAA	AACGCATTTA	ATACTAACTG	1440
	CAGTAGTTCT	TTCAAGAATC	TTTAGAGATA	AGGATTGCAC	ATTGGAAAAG	TAAACCATGT	1500
	TTCATTCCTT	TTTCCCTATT	TATATTGAAA	GAAATAGGCC	AGCAGAGACT	TAGGGATTTT	1560
	AAATTGGCTT	GCTTTTTAGC	TGTTTCAGTC	ACCAGTGAAG	AGCCTATGTG	CATTTTGTAG	1620
10	TAGATAATGT	AAAATTTGTC	ATCTTTTTCT	TTTCTTTTTT	TTAGAATAGC	TGATATTTTG	1680
	ATAACAATCT	CTAATTTGCA	TGGGCACCAC	ATTTCTTATA	TTAAAAGAAT	TAGTGTTTTG	1740
	GCTTCTGTAC	TGCTTATGGT	TGTAGGATTC	AGGGGTTAAT	GGAATCACAG	AAATGATATT	1800
	CTGCAAGAAT	TTCTTTTAAA	TAAAAAGTTT	GGGGGTGCAA	TATAAGAAGT	TTATATAATA	1860
	TGCAGTACAT	TATCCAAAAG	AGAAGGTAGT	TAATGCAGTA	GAAAGTAGTG	GTAATAATTC	1920
15	CTTTTT						

#### SEQ ID NO: 247 PBY4 Protein sequence:

Protein Accession #:

MSDFDSNPFA DPDLNNPFKD PSVTQVTRNV PPGLDEYNPF SDSRTPPPGG VKMPNVPNTQ 60
PAIMKPTEEH PAYTQIAKEH ALAQAELLKR QEELERKAAE LDRREREMQN LSQHGRKNIW 120
PPLPSNFPVG PCFYQEFSVD IPVEFQKTVK LMYYLWMFHA VTLFLNIFGC LAWFCVDSAR 180
AVDFGLSILW FLLFTPCSFV CWYRPLYGAF RSDSSFRFFV FFFVYICQFA VHVLQAAGFH 240
NWGNCGWISS LTGLNQNIPV GIMMIIIAAL FTASAVISLV MFKKVHGLYR TTGASFEKAQ 300
QEFATGVMSN KTVQTAAANA ASTAASSAAQ NAFKGNQI

SEQ ID NO:248 PBH2 DNA sequence

Nucleic Acid Accession#: none found

Coding sequence:

1-613 (underlined sequence corresponds to start and stop codon)

35
ATGAGAGACA ATAAATCGTG TGCTTTTTTC ATGGGAAAGT TAAATGTTTG TTTTGAAGGC 60
ACAGTAATAG CAGGCTATTC AGTGTTTGCC ACTACCTGCA TCATTCATCT GGCTGTAGCC 120
AGTGCACTAC AATTTCCTAA AAAGTCTTCT CACCCTCACA GGACTGCTCT ACACTTAGCC 180
TCTGCCAATG GAAATTCAGA AGTAGTAAAA CTCCTCGCTGG ACAAGACGATG TCAACTTAAT 240
ATCCTTGACA ACAAAAAGAG GACAGCTCTG ACAAAGGCCG TACAATGCCA GGAAGATGAA 300
TGTGCGTTAA TGTTGCTGGA ACATGGCACT GATCCGAATA TTCCAGATGA GTATGGAAAT 360
ACCGCTCTAC ACTATGCTAT CTACAATGAA GATAAATTAA TGGCCAAAGC ACTGCTCTTA 420
TACGGTGCTG ATATCGAATC AAAAAACAAG CATGGCCTCA CACCACTGTT ACTTGGTGTA 480
CATGAGCAAA AACAGCAGT GGTGAAATTT TTAATCAAGA AAAAAGCAAA TTTAAATGCA 540
CTGGATAGAT ATGGAAGGTG TGTGACCTTG GGAACGTTAT TTACCACCAA ATATGTTGTC 600
ATATATGAAA AGTAG

45

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#### SEQ ID NO:249 PBH2 Protein sequence:

Protein Accession #: none found

50 MRDNKSCAFF MGKLNVCFEG TVIAGYSVFA TTCIIHLAVA SALQFPKKSS HPHRTALHLA 60 SANGNSEVVK ILLDRRCQLN ILDNKKRTAL TKAVQCQEDE CALMLLEHGT DPNPDEYGN 120 TALHYAIYNE DKLMAKALLL YGADIESKNK HGLTPLLLGV HEQKQQVVKF LIKKKANLNA 180 LDRYGRCVTL GTLFTTKYVV IYEK

55

SEQ ID NO:250 PBJ1 DNA sequence

Nucleic Acid Accession#: XM_005829

Coding sequence: 1-3043 (underlined sequence corresponds to start and stop codon)

60 ATGGTGATCA TCTATCTTTC TTTCTGCAAT TATTACATGG AGTTCTACAG AGAAGAGCTT 60 CCCCACATTG ACTATTTGAT TGACATTCAG TTTGCAACAG GAAAGGTTAC TCAGCCGGGA 120
GAGGACACTT CCTACCATCA ATGCGCTCAG CTTGAAGCCA GAGACGAAGG CACCGACAGT 180 TTATTATA ACAATGCAG CAGCCCACG CTGAAGACAC GAACGCGCTG TTATGGAACC 240
CCCAGAGGTC TCCCCCATCG TAGCCTGCTC CAGCCGACTC CGCCCACATG TAAAACGAAG 300 ATCAGGAGCA GATTTGAAGA ATTACAAAGT GAATTGGTGC CAGTCAGCAT GTCAGAGACA 360 65 GACCACATAG CCTCTACTTC CTCTGATAAA AATGTTGGGA AAACACCTGA ATTAAAGGAA 420 GACTCATGCA ACTTGTTTTC TGGCAATGAA AGCAGCAAAT TAGAAAATGA GTCCAAACTA 480 TTGTCATTAA ACACTGATAA AACTTTATGT CAACCTAATG AGCATAATAA TCGAATTGAA 540 GCCCAGGAAA ATTATATTCC AGATCATGGT GGAGGTGAGG ATTCTTGTGC CAAAACAGAC 600 70 ACAGGCTCAG AAAATTCTGA ACAAATAGCT AATTTTCCTA GTGGAAATTT TGCTAAACAT 660 ATTTCAAAAA CAAATGAAAC AGAACAGAAA GTAACACAAA TATTGGTGGA ATTAAGGTCA 720 TCTACATTTC CAGAATCAGC TAATGAAAAG ACTTATTCAG AAAGCCCCTA TGATACAGAC 780 TGCACCAAGA AATTTATTTC AAAAATAAAG AGCGTTTCAG CATCAGAGGA TITGTTGGAA 840 GAAATAGAAT CTGAGCTCTT ATCTACGGAG TTTGCAGAAC ATCGAGTACC AAATGGAATG 900 75 AATAAGGGAG AACATGCATT AGTTCTGTTT GAAAAGTGTG TGCAAGATAA ATATTTGCAG 960 CAGGAACATA TCATAAAAAA GTTAATTAAA GAAAATAAGA AGCATCAGGA GCTCTTCGTA 1020 GACATTTGTT CAGAAAAAGA CAATTTAAGA GAAGAACTAA AGAAAAGAAC AGAAACTGAG 1080 AAGCAGCATA TGAACACAAT TAAACAGTTA GAATCAAGAA TAGAAGAACT TAATAAAGAA 1140 GTTAAAGCTT CCAGAGATCA ACTAATAGCT CAAGACGTTA CAGCTAAAAA TGCAGTTCAG 1200

CAGTTACACA AAGAGATGGC CCAACGGATG GAACAGGCCA ACAAGAAATG TGAAGAGGCA 1260 CGCCAAGAAA AAGAAGCAAT GGTAATGAAA TATGTAAGAG GTGAGAAGGA ATCTTTAGAT 1320 CTTCGAAAGG AAAAAGAGAC ACTTGAGAAA AAACTTAGAG ATGCAAATAA GGAACTTGAG 1380 AAAACACTA ACAAAATTAA GCAGCTTTCT CAGGAGAAAG GACGGTTGA GGAACTIGAG AGAAATAAA GCAACTTTCT CAGGAGAAAG GACGGTTGAC 1380
GAAACTAAGG AAGGCGAAAC GACTAGACTC ATCAGAGAAA GAACGGTTGAC CAGCTGTAT 1440
GAAACTAAGG AAGGCGAAAC GACTAGACTC ATCAGAGAAA TAGACAAATT AAAGGAAGAC 1500
ATTAACTCTC ACGTCATCAA AGTAAAGTGG GCACAAAACA AATTAAAAGC TGAAATGGAT 1560
TCACACAAAGG AAAACCAAAGA TAAACTCAAA GAAACACAA CAAAATTAAC ACAAGCAAAG 1620
GAAGAAGCAG ATCAGATACG AAAAACTGT CAGGATATGA TAAAAACATA TCAGGAGTCA 1680
GAAGAAATTA AATCAAATGA GCTTGATGCA AAGCTTAGAG TCACAAAAAG AGAACTTGAA 1740
AAACAAATGC AAGAAAAATC TGACCAGCTA GAGATGCATC ATGCCAAAAT AAAGGAACTA 1800 10 GAAGATCTGA AGAGAACATT TAAGGAGGGT ATGGATGAGT TAAGAACACT GAGAACAAAG 1860 GTGAAATGTC TAGAAGATGA ACGATTAAGA ACAGAAGATG AATTATCAAA ATATAAGGAA 1920 ATTATTAATC GCCAAAAAGC TGAAATTCAG AATTTATTGG ACAAGGTGAA AACTGCAGAT 1980 CAGCTACAGG AGCAGCTTCA AAGAGGTAAG CAAGAAATTG AAAATTTGAA AGAAGAAGTG 2040 15 GAAAGTCTTA ATTCTTTGAT TAATGACCTA CAAAAAGACA TCGAAGGCAG TAGGAAAAGA 2100 GAATCTGAGC TGCTGCTGTT TACAGAAAGG CTCACTAGTA AGAATGCACA GCTTCAGTCT 2160 GAATCCAATT CTTTGCAGTC ACAATTTGAT AAAGTTTCCT GTAGTGAAAG TCAGTTACAA 2220 AGCCAGTGTG AACAAATGAA ACAGACAAAT ATTAATTTGG AAAGTAGGTT GTTGAAAGAG 2280 GAAGAACTGC GAAAAGAGGA AGTOCAAACT CTGCAAGCTG AACTGCTTG TAGACAAACA 2340
GAAGTAAAG CATTGAGTAC CCAGGTAGAA GAATTAAAAG ATGAGTTAGT AACTCAGAGA 2400
CGTAAACATG CCTCTAGTAT CAAGGATCTC ACCAAACAAC TTCAGCAAGC ACGAAGAAAA 2460
TTAGATCAGG TTGAGAGTGG AAGCTATGAC AAAGAAGTCA GCAGCATGGG AAGTCGTTCT 2520 20 AGTTCATCAG GGTCCCTGAA TGCTCGAAGC AGTGCAGAAG ATCGATCTCC AGAAAATACT 2580 GGGTCCTCAG TAGCTGTGGA TAACTTTCCA CAAGTAGATA AGGCCATGTT GATTGAGAGA 2640 25 ATAGTTAGGC TGCAAAAAGC ACATGCCCGG AAAAATGAAA AGATAGAATT TATGGAGGAC 2700 CACATCAAAC AACTGGTGGA AGAAATTAGG AAAAAAACAA AAATAATTCA AAGTTATATT 2760 TTACGAGAAG AATCAGGCAC ACTITCTTCA GAGGCATCTG ATTTTAACAA AGTTCATTTA 2820 AGTAGACGGG GTGGCATCAT GGCATCTTTA TATACATCCC ATCCAGCTGA CAATGGATTA 2880 ACATTGGAGC TCTCTTTGGA AATCAACCGA AAATTACAGG CTGTTTTGGA GGATACGTTA 2940 ACATIGGAGC TCICTITIGA AAICAACCGA AAATTACAGG CTGITTIGGA GGATACGITA 2940
CTAAAAAATA TTACTTTGAA GGAAAATCTA CAAACACTTG GAACAGAAAT AGAACGTCTT 3000
ATTAAACACC AGCATGAACT AGAACAGAGG ACAAAGAAAA CC<u>TAA</u>AACAA GCCTCTTGCT 3060
CAGTAAAGAG ACAAAAGCCA CACAGGAGTA GGTGCCACTG ACCTCTATTG TTGGAGACTT 3120
TGTTCCACTT TITGTTTCAG CCAGTAAAAA TATTGTTTTG CTTCATCTGT ACACAAAAAA 3180
ATACCCTTTT ACAATATGAA TGCATTGCTG TATATACTGT AAGACTGAAA GCTTTGATGA 3240
AATTTGTTTT TGTATGGTGC AATATGACAG CCTGTCATTG AATCTAAACA ACTTAATTTG 3300 30 35 CTIGTATTCA TAAGAAGTGT TGAACATTAC AAGGGCTTTT AT

# 40 Protein Accession #: SEQ ID NO:251 PBJ1 Protein sequence; NP_060487

MVIIYLSFCN YYMEFYREEL PHIDYLIDIQ FATGKVTQPG EDTSYHQCAQ LEARDEGTDS 60
ILLINGSSAT LKTRTRCYGT PRGLPHRSILL QPTPPTCKTK IRSRFEELQS ELVPVSMSET 120
DHIASTSSDK NVGKTPELKE DSCNLFSGNE SSKLENESKL LSLNTDKTLC QPNEHNNRIE 180
AQENYIPDHG GGEDSCAKTD TGSENSEQIA NFPSGNFAKH ISKTNETEGK VTQILVELRS 240
STFPESANEK TYSESPYDTD CTKKFISKIK SVSASEDLLE EIESELLSTE FAEHRVPNGM 300
NKGEHALVIF EKCVQDKYLQ QEHIIKKLIK ENKKHQELFV DICSEKDNILR EELKKRTETE 360
KQHMNTIKQL ESRIEELNKE VKASRDQLIA QDVTAKNAVQ QLHKEMAQRM EQANKKCEEA 420
RQEKEAMYMK YVRGEKESLD LRKEKETLEK KLRDANKELE KNTNKIKQLS QEKGRLHQLY 480
ETKEGETTRL IREIDKLKED INSHVIKVKW AQNKLKAEMD SHKETKDKLK ETTTKLTQAK 540
EEADQIRKNC QDMIKTYQES EEIKSNELDA KLRVTKGELE KQMQEKSDQL EMHHAKIKEL 600
EDLKRTFKEG MDELRTLRTK VKCLEDERLR TEDELSKYKE IINRQKAEIQ NILDKVKTAD 660
QLQEQLQRGK QEIENLKEEV ESLNSLINDL QKDIEGSRKR ESELLLFTER LTSKNAQLQS 720
ESNSLQSQFD KVSCSESQLQ SQCEQMKQTN INLESRILKE EEIRKEEVQT LQAELACRQT 780
EVKALSTQVE ELKDELVTQR RKHASSIKDL TKQLQQARRK LDQVESGSYD KEVSSMGSRS 840
SSSGSLNARS SAEDRSPENT GSSVAVDNFP QVDKAMLIER IVRLQKAHAR KNEKIEFMED 900
HIKQLVEEIR KKTKIIQSYI LREESGTILSS EASDFNKVHL SRRGGIMASL YTSHPADNGL 960
TLELSLEINR KLQAVLEDIL LKNITLKENL QTLGTEIERL IKHQHELEQR TKKT

SEQ ID NO:252 PBJ6 DNA sequence Nucleic Acid Accession#: D83760 Coding sequence: 56-1459

60

56-1459 (underlined sequence corresponds to start and stop codon)

65	1	11	21	31	41	51	
-	ī	ī	ĩ	Ĩ	ï	Ĭ	
	TTGCCGTGAA	GGGCTGTGCG	GTTCCCGTGC	GCGCCGGAGC	CTGCTGTGGC	CTCTTATGCA	60
	CTCCACCACC	CCCATCAGCT	CCCTCTTCTC	CTTCACCAGC	CCCGCAGTGA	AGAGACTGCT	120
	AGGCTGGAAG	CAAGGAGATG	AAGAGGAAAA	GTGGGCAGAG	AAGGCAGTGG	ACTCTCTAGT	180
70	Gaagaagtta	AAGAAGAAGA	AGGGAGCCAT	GGACGAGCTG	GAGAGGGCTC	TCAGCTGCCC	240
	GGGGCAGCCC	AGCAAATGCG	TCACGATTCC	CCGCTCCCTG	GACGGGCGGC	TGCAGGTGTC	300
	CCACCGCAAG	GGCCTGCCCC	ATGTGATTTA	CTGTCGCGTG	TGGCGCTGGC	CGGATCTGCA	360
	GTCCCACCAC	GAGCTGAAGC	CGCTGGAGTG	CTGTGAGTTC	CCATTTGGCT	CCAAGCAGAA	420
~~	AGAAGTGTGC	ATTAACCCTT	ACCACTACCG	CCGGGTGGAG	ACTCCAGTAC	TGCCTCCTGT	480
75	GCTCGTGCCA	AGACACAGTG	AATATAACCC	CCAGCTCAGC	CTCCTGGCCA	AGTTCCGCAG	540
	CGCCTCCCTG	CACAGTGAGC	CACTCATGCC	ACACAACGCC	ACCTATCCTG	ACTOTTTCCA	600
	GCAGCCTCCG	TGCTCTGCAC	TCCCTCCCTC	ACCCAGCCAC	GCGTTCTCCC	AGTCCCCGTG	660
	CACGGCCAGC	TACCCTCACT	CCCCAGGAAG	TCCTTCTGAG	CCAGAGAGTC	CCTATCAACA	720
	CTCACTTGAC	ACACCACCCC	ጥርርምምልጥናል	TYCCCACAGAA	COCTOTORAGA	CCCAGAGTGG	780

	CCAACCTGTA	GATGCCACAG	CTGATAGACA	ТСТАСТССТА	TOGATACCAA	ATGGAGACTT	840
			AGCCCCAGCA				900
							960
			TCCAGGCTTC				
. 5						TAAACAGAAA	1020
	CTCAACGATA	GAAAATACCA	GGAGACATAT	aggaaagggt	GTGCACTTGT	ACTACGTCGG	1080
			GCGTGAGTGA				1140
	CAACTATCAA	CACGGCTTCC	ACCCAGCTAC	CGTCTGCAAG	ATCCCCAGCG	GCTGCAGCCT	1200
			TCTTCGCTCA				1260
			CCAAGATGTG				1320
10			AGGATGTCAC				1380
10							
			TGGACAAAGT				1440
	CATTTCTTCA	GTGTCTTAAC	AGTCATGTCT	TAAGCTGCAT	TTCCATAGGA	T	
1 ~							
15		SEQ II	D NO:253 P <u>BJ6 P</u> r	otein sequence:			
	Protein Accessio	n#: NP_00	5896 ·				
		_					
	MHSTTPISSI.	FSFTSPAVKR	LLGWKOGDE	EKWAKKAVI	OS LVKKI KKI	KG AMDELER	ALS 60
						PL ECCEFPFGS	
20							
20						LMPHNATYPD	
						HATEASETQ	
	SGQPVDATA	D RHVVLSIPN	G DFRPVCYEE	P QHWCSVAY	<b>YE LNNRVGE</b>	IFQ ASSRSVLII	DG 300
	FTDPSNNRNI	R PCLGLLSNV	N RNSTIENTRE	RHIGKGVHLY	Y VGGEVYAE	CV SDSSIFVQS	R 360
	NCNYOHGFH	P ATVCKIPSG	C SLKVFNNOL	FAOLLAOSVI	IH GFEVVYEL	TK MCTIRMSF	VK 420
25			EI HLHGPLOW				
	CEO ID NO-254	PBJ8 DNA sequer	300			•	
30	Nucleic Acid Acc					4	
30	Coding sequence	3: 4/2-4	377 (underlined se	quence correspon	as to start and sto	p coaon)	
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~ ~	TGCAGGTTTG	CAGGGTCTGA	GATTACTTGG	GCTTTTCCTG	CCTTTTTCTT	TTGCTTAAGG	60
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	TGGGGACACT	TGGTTGATGC	AGTCTCTCTC	TCTCTTTCTC	GGTGTTTATA	ACAAAACAAA	180
			TTGTAATGGT				240
			AGGGTATGTT				300
			TCTGACTATT				360
40			TTGCTGAGAA				420
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			TCAAGGTTCA				480
			CTTTGATGAC				540
			TGAGTCTGGA				600
45			CTCCCACGCA				660
43			CATTGACTCT				720
			TAATGGGTTT				780
			GAAAGGAGAT				840
			CCCGATCTCC				900
50	attgaggtgg	ATGACCCCCC	TGACAAGGAG	GACATGCGAT	CAAGCTTCAG	GTCGAATGTG	960
50	TTGACGGGGT	CGGCTCCCCA	GCAGGACTAC	GATAAGCTGA	AGGCACTCGG	AGGGGAAAAC	1020
	TCCAGCAAAA	CTGGACTCTC	TACGTCAGGC	AATGTGGAGA	AAAACAAAGC	TGTTAAGAGA	1080
·	GAAACAGAAG	CCAGTTCTAT	AAACCTGAGT	GTTTATGAAC	CTTTTAAAGT	CAGAAAAGCA	1140
	GAGGATAAAT	TGAAGGAAAG	CTCTGACAAG	GTGCTGGAAA	ACAGAGTCCT	AGATGGGAAG	1200
			CACCAGCCTC				1260
55			CATCGCTGCC				1320
			AGTGGCCAAT				1380
			CGCTGACAAG				1440
			GCAACCGGAT				1500
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00						ATTGCCAGAA	
						GCCTCTGTG	
						CAGGGCGCCT	1740
						CAAACAGGTC	1800
C 5						GGCAGGATCC	
65	CAAGTCATTA	ATTTGAAGCT	CGCTAACAAC	ACCACGGTGA	AAGCCACGGT	CATATCTGCT	1920
	GCCTCTGTCC	AGAGTGCCAG	CAGCGCCATC	ATTAAAGCTG	CCAACGCCAT	CCAGCAGCAA	1980
	ACTGTCGTGG	TGCCGGCATC	CAGCCTGGCC	AATGCCAAAC	TCGTGCCAAA	GACTGTGCAC	2040
	CTTGCCAACC	TTAACCTTTT	GCCTCAGGGT	GCCCAGGCCA	CCTCTGAACT	CCGCCAAGTG	2100
			AATAAAGCAG				2160
70			CCAGGTGGTG				2220
						CCTCAGTCCT	
						GTGTGGGGAC	
	CCCCCCCCCCCC		THE CARE				
			montos cocas o				
	TCCTTTGCAC	TTGAAAAGAG	TCTGACCCAG				
75	TCCTTTGCAC GTAACGTGCA	TTGAAAAGAG ACCATTGTAC	AAAGAACCTC	GTTTTTTACA	ACAAATGCAG	CCTCCTTTCC	2460
75	TCCTTTGCAC GTAACGTGCA CATGCCCGTG	TTGAAAAGAG ACCATTGTAC GGCATAAGGA	AAAGAACCTC GAAAGGGGTG	GTTTTTTACA GTAATGCAAT	ACAAATGCAG GCTCCCACTT	CCTCCTTTCC AATTTTAAAG	2460 2520
75	TCCTTTGCAC GTAACGTGCA CATGCCCGTG CCAGTCCCAG	TTGAAAAGAG ACCATTGTAC GGCATAAGGA CAGATCAAAT	AAAGAACCTC GAAAGGGGTG GATAGTTTCT	GTTTTTTACA GTAATGCAAT CCGTCAAGCA	ACAAATGCAG GCTCCCACTT ATACTTCCAC	CCTCCTTTCC AATTTTAAAG TTCAACTTCC	2460 2520 2580
75	TCCTTTGCAC GTAACGTGCA CATGCCCGTG CCAGTCCCAG ACTCTTCAGA	TTGAAAAGAG ACCATTGTAC GGCATAAGGA CAGATCAAAT GCCCTGTGGG	AAAGAACCTC GAAAGGGGTG GATAGTTTCT AGCTGGCACA	GTTTTTTACA GTAATGCAAT CCGTCAAGCA CACACTGTCA	ACAAATGCAG GCTCCCACTT ATACTTCCAC CAAAAATTCA	CCTCCTTTCC AATTTTAAAG TTCAACTTCC GTCTGGCATA	2460 2520 2580 2640
75	TCCTTTGCAC GTAACGTGCA CATGCCCGTG CCAGTCCCAG ACTCTTCAGA ACTGGGACAG	TTGAAAAGAG ACCATTGTAC GGCATAAGGA CAGATCAAAT GCCCTGTGGG TCATATCGGC	AAAGAACCTC GAAAGGGGTG GATAGTTTCT AGCTGGCACA TCCTTCAAGC	GTTTTTTACA GTAATGCAAT CCGTCAAGCA CACACTGTCA ACTCCCATCA	ACAAATGCAG GCTCCCACTT ATACTTCCAC CAAAAATTCA CCCCAGCCAT	CCTCCTTTCC AATTTTAAAG TTCAACTTCC GTCTGGCATA GCCCCTAGAT	2460 2520 2580 2640 2700
	TCCTTTGCAC GTAACGTGCA CATGCCCGTG CCAGTCCCAG ACTCTTCAGA ACTGGGACAG	TTGAAAAGAG ACCATTGTAC GGCATAAGGA CAGATCAAAT GCCCTGTGGG TCATATCGGC	AAAGAACCTC GAAAGGGGTG GATAGTTTCT AGCTGGCACA TCCTTCAAGC	GTTTTTTACA GTAATGCAAT CCGTCAAGCA CACACTGTCA ACTCCCATCA	ACAAATGCAG GCTCCCACTT ATACTTCCAC CAAAAATTCA CCCCAGCCAT	CCTCCTTTCC AATTTTAAAG TTCAACTTCC GTCTGGCATA	2460 2520 2580 2640 2700
75 80	TCCTTTGCAC GTAACGTGCA CATGCCCGTG CCAGTCCCAG ACTCTTCAGA ACTGGGACAG GAAGACCCCT	TTGAAAAGAG ACCATTGTAC GGCATAAGGA CAGATCAAAT GCCCTGTGGG TCATATCGGC CCAAACTGTG	AAAGAACCTC GAAAGGGGTG GATAGTTTCT AGCTGGCACA TCCTTCAAGC TAGACATAGT	GTTTTTTACA GTAATGCAAT CCGTCAAGCA CACACTGTCA ACTCCCATCA CTAAAATGTT	ACAAATGCAG GCTCCCACTT ATACTTCCAC CAAAAATTCA CCCCAGCCAT TGGAGTGTAA	CCTCCTTTCC AATTTTAAAG TTCAACTTCC GTCTGGCATA GCCCCTAGAT	2460 2520 2580 2640 2700 2760

PCT/US01/32045 WO 02/30268

	ACTTGCACTA	TCTGCCAGAT	GCTGCTTCCT	AACCAGTGCA	GTTATGCATC	ACACCAGAGA	2880
		ACAAATCTCC					2940
		CCCACGTCAC					3000
		GCAATGTTGT					3060
5		AAGTCTTCTA					3120
•		CCCACGCCTA		_			3180
		GTTCCATGTG					3240
		ACATTGAAAA					3300
		AGAAGCAACT					3360
10		GGCCTCCAAA					3420
		CAAATCAGAA					3480
		CTCCATCTCC					3540
		GTTGGGAGTG					3600
		AGCACGGGAA					3660
15		CCCACAGCCT					3720
10		GCTCGCACTG					3780
		TCCAGCTGAT					3840
		AGGAGGAAAC					3900
		AACCAGTTCT					3960
20		AAATCAATGT					4020
20		TGCAATTCCA					4080
		AGTGTGGCCT					4140
		AGTTAAAGGA					4200
		AGAACAAACC					4260
25		AAGTGTGCGC					4320
~~		GCATGGCCTT					4380
		ATGAGGAAAA					4440
		TATAATAGAG					4500
		CCTTCACCTC					4560
30		TTTGTATATA					4620
<b>J</b>		TAGTGGAAAA					4680
		AAACAGAGTT					4740
		GGATTTTGAA					4800
		TTTTAGAATT					4860
35		TAAGTGTCTT					4920
		AACTGCACTC					4980
		AGTCTTGCAG					5040
•		GGAATGCTGA					5100
		ATGGGATTTG					5160
40	GGAGGGAAAG	TAACGAAAGG	GCTGGACTAC	TATAAAAGTT	ACAAATACGT	AGTTAGACCA	5220
		ATAGTCAGGT					5280
	AGCTAAGAAT	ATCAAGTATT	TCTCTGGCTC	TTGACAGAAA	AAAATCAGTT	GACTTAACCC	5340
	TTTGCTGTCA	AAAGAGTTGG	CGTTTCCTGT	TCTGGGTGCT	ACTGCCAAAC	GTTATGGTAC	5400
	TTAGAGTCGG	GATGCACAAC	TTCAACCACC	GACTTATCAA	TGCAGCCGCC	TGTGTATTGC	5460
45	AATTGGCCGT	TACCTTAAGC	ACTGAGCCAC	CCGGGTTTAG	TTCAGCCATT	TCAAGAAGTA	5520
	TATTTAACGT	CGGTAGTTCT	GCTTTATTAA	AATGCAGCAG	AGGTACTCTT	CTGTCCCTTC	5580
	CGTTTATAGT	TCTCTGAGAG	AGTTCTATTT	TTTGGTTTTG	TTTTGTGTTT	TCTTTTGCAT	5640
	TTTGTATCTT	GTATTTATCC	CTGAACATGT	TTTGTACCTT	TTTTTTTTT	TTTTTTTTAA	5700
~^	GAAAAGGAAT	TCTTTTGTGT	ATATATAGAT	ACTTGCATGA	TATACTGTAG	TCAATGTTCG	5760
50	GTTCCTCAAA	AGGTCTTGCT	GCTGTCAGGT	GTTATGCACT	CCATCCATCA	TAACTGTATG	5820
	AAACACATTT	CATATGTAAA	TAAACGTGGG	ACATTTG			

#### SEQ ID NO:255 PBJ8 Protein sequence; 55

Protein Accession #:

MKTPDFDDLL AAFDIPDMVD PKAAIESGHD DHESHMKQNA HGEDDSHAPS SSDVGVSVIV 60 KNVRNIDSSE GGEKDGHNPT GNGLHNGFLT ASSLDSYSKD GAKSLKGDVP ASEVTLKDST 120 FSQFSPISSA EEFDDDEKIE VDDPPDKEDM RSSFRSNVLT GSAPQQDYDK LKALGGENSS 180 60 KTGLSTSGNV EKNKAVKRET EASSINLSVY EPFKVRKAED KLKESSDKVL ENRVLDGKLS 240 SEKNDTSLPS VAPSKTKSSS KLSSCIAAIA ALSAKKAASD SCKEPVANSR ESSPLPKEVN 300 DSPRAADKSP ESQNLIDGTK KPSLKQPDSP RSISSENSSK GSPSSPAGST PAIPKVRIKT 360 DSPRADKSP ESQNLIDGTK KPSLKQPDSP RSISSENSSK GSPSSPAGST PAIPKVRIKT 360

KTSSGEIKR TVTRVLPEVD LDSGKKPSEQ TASVMASVTS LLSSPASADY LSSPPRAPLQ 420

SAVVTNAVSP AELTPKQVTI KPVATAFLPV SAVKTAGSQV INLKLANNTT VKATVISAAS 480

VQSASSAIIK AANAIQQQTV VVPASSLANA KLVPKTVHLA NLNILPQGAQ ATSELRQVLT 540

KPQQQIKQAI INAAASQPPK KVSRVQVVSS LQSSVVEAFN KVLSSVNPVP VYIENLSPPA 600

NAGITLPTRG YKCLECGDSF ALEKSLTQHY DRRSVRIEVT CNHCTKNLVF YNKCSLLSHA 660

RGHKEKGVVM QCSHLILKPV PADQMIVSPS SNTSTSTSTL QSPVGAGTHT VTKIQSGITG 720

TVISAPSSTP ITPAMPLDED PSKLCRHSLK CLECNEVFQD ETSLATHFQQ AADTSQKTC 780

TICQMLLPNQ CSYASHQRIH QHKSPYTCPE CGAICRSVHF QTHVTKNCLH YTRVGFRCV 840

HCNVLVYSDVA AI KSHIGGSH CEVEYKCPIC PMARKSAPST HSHAYTOHPG IKIGEPKITY 900 65 70 HCNVVYSDVA ALKSHIQGSH CEVFYKCPIC PMAFKSAPST HSHAYTQHPG IKIGEPKIIY 900 KCSMCDTVFT LQTLLYRHFD QHIENQKVSV FKCPDCSLLY AQKQLMMDHI KSMHGTLKSI 960 EGPPNLGINL PLSIKPATON SANONKEDTK SMNGKEKLEK KSPSPVKKSM ETKKVASPGW 1020 TCWECDCLFM QRDVYISHVR KEHGKQMKKH PCRQCDKSFS SSHSLCRHNR IKHKGIRKVY 1080
ACSHCPDSRR TFTKRLMLEK HVQLMHGIKD PDLKEMTDAT NEEBTEIKED TKVPSPKRKL 1140
EEPVLEFRPP RGAITQPLKK LKINVFKVHK CAVCGFTTEN LLQFHEHIPQ HKSDGSSYQC 1200
RECGLCYTSH VSLSRHLFIV HKLKEPQPVS KQNGAGEDNQ QENKPSHEDE SPDGAVSDRK 1260 75 CKVCAKTFET EAALNTHMRT HGMAFIKSKR MSSAEK

80

#### SEQ ID NO:256 PBM1 DNA sequence Nucleic Acid Accession#: AF111847

Coding sequence: 58-1608 (underlined sequence corresponds to start and stop codon)

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•				ATCTTCAAGC			120
10				AATCCCAGCT			180
10				CACCGGTCAC			240
				TCATGGTTTC			300
				CATCAACATG			360
				CTCTATAGGG			420
15				CTGTGGCTTG			480
13				TTTGCCTCTC			540
				CCATCTTCTT			600
				GAGCAAGGAC			660
				TCTATCATAA			720
20				TTGGGAGCTC			780
20				GCGGATAAAA			840
				GTTTCATCAT			900
				ATGAACATTA			960
	TCAGACAGAC	TCGGCATGGG	ATTTGGAAAT	TGCAGAAGTG	TTATTTCACA	TTCAGTGACT	1020
05				CCCATTATGG			1080
25	AATGATGACA	GTGACGATTC	ATATTTTACT	TCCAGCTCAA	GTTACTTTGA	CGAGCCAGTG	1140
	GAGTTAAGGA	GCAGTTCTTT	CTCTAGCTGG	GATGACAGTT	CAGATTCCTA	TTGGAAAAA .	1200
	GAGACCAGCA	AAGATACTGA	AACAGTTCTG	AAAACCACAG	<b>GCTATTCAGA</b>	CAGACCTACT	1260
	GCTCGCCGCA	AGCCAGATTA	TGAGCCAGTT	GAAAATACAG	ATGAGGCCCA	GAAGAAGTTT	1320
20				TATTTTGGAA			1380
30	GAGACCAGGG	CCCGCCTAGA	GAGGCTGTCG	GCAAGTTCCT	CCATAAGCTC	GGCTGATCTG	1440
	TTCGAGGAGC	CGAGGAAGCA	GCCAGCAGGG	AACTACAGCC	TGTCCAGTGT	GCTGCCCAAC	1500
				GTGAGATCGG			1560
				GATCGCTACG			1620
25	GATGTGTATT	TCCTGGAGAA	ATTCCTCTTT	AAATGAACAA	GTAACCACAT	CTCAGGCGGC	1680
35	AGTGAAGTCC	AGATAGTTTT	GCAGATTGTT	TIGCTACTIT	TTCATATGGT	ATATGTTTCT	1740
	GATTTTTAAT	ATTTCTTTTG	AGAAATTCTG	AGTTCTGATG	TAGGAGCTTT	CCTGTGATTT	1800
	CTGTTTCACG	TTCCTTCCTG	TCACACCCTC	CTTTGGCGTC	TCTGTGTATA	TCCTTGCTTT	1860
	ATTTTCTTGG	AACCTTTGAT	TTCAACACTG	AGGGCCTGGA	GACCTCGGCT	CCTCCTGCTC	1920
40	CTGAACCAGG	AGGCTTCATG	TGGGGGAGGA	GGAGAGGTCT	CCATGTGACA	CATGGGCTCA	1980
40	GGGCTGCCAG	AATCAGCGGA	TGCTGGATGG	GCCTGCAGAA	ACAACACTCA	CCACACACAC	2040
	TTCCTTCAAA	AGACCAAAAG	TGACTGGTGT	CTCGTGTGAC	AGATTGCTTC	ATTTATGTTT	2100
	CTACATAGTA	AGGTGACTGC	CAAATAATAT	TTGAAGTCAT	CTGTCTCTTT	GTAAATTATT	2160
	TTATATGACC	ATTTAAATTA	AAAATGTTTT	TCAGTGAGTG	CTTTTAACAA	ACTTAAGCTT	2220
. ~	CTGCCCTGCC	AAGGGAATTA	ATGTTATCTT	GTGAAAGGTG	TIGCTGTTIG	AATTGATGAG	2280
45	AAATGGAAGA	TGAGAACTCC	CTAAGAGTTC	TCATAATAAA	TCATCTCATC	ACAAATCAAT	2340
	ACGGTATACA	GAGTTAAAGT	GGAATGAGGT	AAGAAGATAC	AGCTACAGAA	AATAGTTGCG	2400
	TGTATGGGAG	AACAGTCATT	GTAATTGGGT	AGTTTTGTTA	TTTATAATTA	TTAAATCTTG	2460
	CTTTTCAGAA	ATTACCGAAT	GTGTATAAAC	AAATAAAGAA	AAATAATTTA	GCTGTGTTTT	2520
	AGACAGCATT	AGAATATATT	GTTCAGCACA	GTAAAATATA	TTTGAAATTT	GATAAGCCAA	2580
50	AAATGTGGTT	TTGAATGAAT	ATTTTGTGAA	TCTTTCTTAA	AAGCTCAAAT	TIGTAGACTT	2640
	CTAAATAGAA	TAAACACTTG	CAGCAGAAAA	AAAAAAAAA	АААААААА	AAAAAAAAA	2700
				ААААААААА			2760
	AAAAAAAAA						
55			•				

55
SEQ ID NO:257 PBM1 Protein sequence:
PBM1 Protein sequence: CAB76901

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FIRSTELDSN WSWFQLRCMQ VGGNASASSF FHQHGCSTND TNAKYNSRAA QLYREKIKSL 120
ASQATRKHGT DLWLDSCVVP PLSPPPKEED FFASHVSPEV SDTAWASAIA EPSSLTSRPV 180
ETTLENNEGG QEQFSVEGL NVPTKATLEV SSIIKKKPNQ AKKGLGAKKG SLGAQKLANT 240
CFNEIEKQAQ AADKMKEQED LAKVVSKEES IVSSLRLAYK DLEIQMKKDE KMNISGKKNV 300
DSDRLGMGFG NCRSVISHSV TSDMQTTEQE SPIMAKPRKK YNDDSDDSYF TSSSSYFDEP 360
VELRSSSFSS WDDSSDSYWK KETSKDTETV LKTTGYSDRP TARRKPDYEP VENTDEAQKK 420
FGNVKAISSD MYFGRQSQAD YETRARLERL SASSSISSAD LFEEPRKQPA GNYSLSSVLP 480
NAPDMAQFKQ GVRSVAGKLS VFANGVVTSI QDRYGS

70 <u>SEQ ID NO:258 PBM4 DNA sequence</u> Nucleic Acid Accession#: D30891

Coding sequence: 1-4032 (underlined sequence corresponds to start and stop codon)

75 ATGATACTG TCATGAAGCA GACACATGCT GACACACCTG TTGATCATTG TCTATCTGGC 60
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CTTGAAATGC AGAATCCAAA TTTGAACAAT AAAAGAATGTT GTTTTCACCTTTACGTTGAAT 180
GGAAACTCCA GAAAATTAGA CCGTAGTGTG TTTTACAGCAT ATGGTAAACC CAGCGAGAGT 240
ATCTACTCAG CCCTGAGTGC TAATGACTAT TTCAGTGAAA GGATAAAGAA TCAGTTTAAT 300
AAGAACATTA TTGTTTATGA AGAAAAGACA ATAGATGGAC ATATAAATTT AGGAATGCCT 360
CTCAAGTGCC TGCCTAGTGA TTCTCATTTT AAAATTACAT TTGGTCAAAG AAAGAGTAGC 420

AAAGAAGATG GACACATATT ACGCCAATGT GAAAATCCAA ACATGGAATG CATTCTTTTT 480 CATGTTGTTG CTATAGGAAG GACAAGAAAG AAGATTGTTA AGATCAACGA ACTTCATGAA 540 AAAGGAAGTA AACTTTGTAT TTATGCCTTG AAGGGTGAGA CTATTGAAGG AGCCTTATGC 600 AAGGATGGCC GTTTTCGGTC TGACATAGGT GAATTTGAAT GGAAACTAAA GGAAGGTCAT 660
AAGAAAATTT ATGGAAAACA GTCCATGGTG GATGAAGTAT CTGGAAAAGT 720
GACATTTCAA AAAAAAAACC ATTACAACAG AAAGATATCC ATAAAAAAAT TAAACAGAAT 780
GAAAGTGCCA CTGATGAAAT TAATCACCAG AGTCTGATAC AGTCTAAGAA AAAAGTCCAC 840
AAACCAAAGA AAGATGGAGA GACCAAAGAT GTAGAACACA GCAGAGAGCA AATTCTCCCA 900 CCTCAGGATC TAAGCCATTA TATTAAAGAT AAAACTCGCC AGACAATTCC CAGGATTAGA 960 10 AATTATTACT TITGTAGTTT GCCCCGAAAA TATAGGCAAA TAAACTCACA AGTTAGACGG 1020 AGGCCGCATC TGGGTAGGCG GTATGCTATT AATCTGGATG TCCAAAAGGA GGCAATTAAT 1080 CTCTTAAAGA ATTATCAAAC GTTGAATGAA GCCATAATGC ATCAGTATCC GAATTTTAAA 1140 GAGGAGGCAC AGTGGGTAAG AAAATATTTT CGGGAAGAAC AAAAGAGAAT GAATCTTTCA 1200 CCAGCTAAGC AATTCAACAT ATATAAAAAG GACTTCGGAA AAATGACTGC AAATTCTGTT 1260 15 TCAGTTGCAA CCTGCGAACA GCTTACATAT TATAGCAAGT CAGTTGGGTT CATGCAATGG 1320 GACAATAATG GAAACACAGG TAATGCTACT TGCTTTGTCT TCAATGGTGG TTATATTTTC 1380 ACCTGTCGAC ATGTTGTACA TCTTATGGTG GGTAAAAACA CACATCCAAG TTTGTGGCCA 1440 GATATAATTA GCAAATGTGC GAAGGTAACC TTCACTTATA CAGAGTTCTG CCCTACTCCT 1500 GACAATTGGT TTTCCATTGA GCCATGGCTT AAAGTGTCCA ATGAAAATCT AGATTATGCC 1560 20 ATTITAAAAC TAAAAGAAAA TGGAAATGCG TITCCTCCAG GACTATGGCG ACAGATTTCT 1620 CCTCAACCAT CTACTGGTTT GATTTATTTA ATTGGTCATC CTGAAGGCCA GATCAAGAAA 1680 ATAGATGGTT GTACTGTGAT TCCTCTAAAC GAACGATTGA AAAAATATCC AAACGATTGT 1740 CAAGATGGGT TGGTAGATCT CTATGATACC ACCAGTAATG TATACTGTAT GTTTACCCAA 1800 AGAAGTTTCC TATCAGAGGT TTGGAACACA CACACGCTTA GTTATGATAC TTGTTTCTCT 1860 25 GATGGGTCCT CAGGCTCCCC AGTGTTTAAT GCATCTGGCA AATTGGTTGC TTTGCATACC 1920 TTTGGGCTTT TTTATCAACG AGGATTTAAT GTGCATGCCC TTATTGAATT TGGTTATTCT 1980 ATGGATTCTA TTCTTTGTGA TATTAAAAAG ACAAATGAGA GCTTGTATAA ATCATTAAAT 2040 GATGAGAAAC TTGAGACCTA CGATGAAGAG AAAGCCCGGC CCAGGCCAGC CTACCGGCGA 2100 CTAGGATGCT TTCGCTTTCG CTCTCGCTTT CCAATACTCG GGACTGGGGA AACCGGGAGA 2160 ATAGAAGCAG GCAAGGACCG CCGTGGGCAC GGGGTCAGTG AGACAGGGTC CTGCTCGCGG 2220 30 CGTCAAGGAG GAGCGCTGTG GGTGTCCCCA GCGCAGCCAA TCGGCTTCCG AAGTAGCTGG 2280 AGCTCTGGAG CCTTTGCTTC CTCAAATACG AGCGGGAACT GCGTTGAGCG CTGGATTCCA 2340 GGCCGAGTGC TGGCGAGGCG CGCAGTCTCT AAAGAGCAAC AGAATAATTG CAGTACTTCT 2400 CTAATGAGGA TGGAGTCTAG AGGAGACCCA AGAGCCACAA CTAATACCCA GGCTCAAAGA 2460 35 TTCCATTCAC CTAAGAAAAA TCCAGAAGAC CAGACCATGC CCCAAAATAG GACAATATAT 2520 GTTACCTTGA AGGCTGTCAG AAAAGAGATA GAAACTCACC AAGGCCAAGA AATGCTTGTG 2580 CGTGGCACAG AAGGAATCAA AGAGTACATA AACCTTGGAA TGCCCCTCAG TTGTTTCCCT 2640 GAAGGTGGCC AGGTGGTCAT TACATTTTCC CAAAGTAAAA GTAAGCAGAA GGAAGATAAC 2700 CACATATTTG GCAGGCAGGA CAAAGCATCG ACTGAATGTG TCAAATTTTA CATTCATGCA 2760 40 ATTGGAATTG GGAAGTGTAA AAGAAGGATT GTTAAATGTG GGAAGCTTCA CAAAAAGGGG 2820 CGCAAACTCT GTGTTTATGC TTTCAAAGGA GAAACCATCA AGGATGCACT GTGCAAGGAT 2880 GGCAGATITIC TITICCITTICT GGAGAATGAT GATTGGAAAC TCATTGAAAA CAATGACACC 2940 ATTTTAGAAA GCACCCAGCC AGTTGATGAA TTAGAAGGCA GATACTTTCA GGTTGAGGTT 3000 GAGAAAAGAA TGGTCCCCAG TGCAGCAGCT TCTCAGAATC CTGAGTCAGA GAAAAGAAAC 3060 45 ACCTGTGTGT TGAGAGAACA AATCGTGGCT CAGTACCCCA GTTTGAAAAG AGAAAGTGAA 3120 AAAATCATTG AAAACTTCAA GAAAAAATG AAAGTAAAAA ATGGGGAAAC ATTATTTGAA 3180
TTGCATAGAA CAACGTTTGG GAAAGTAACA AAAAATTCTT CTTCGATTAA AGTAGTGAAA 3240
CTTCTTGTAC GTCTCAGTGA CTCAGTTGGG TACTTATTCT GGGACAGTGC AACTACGGGT 3300
TACGCCACCT GCTTTGTTTT TAAAGGATTG TTCATTTTAA CTTGTCGGCA TGTAATAGAT 3360 50 AGCATTGTGG GAGACGGAAT AGAGCCAAGT AAGTGGGCAA CCATAATTGG TCAATGTGTA 3420 AGGGTGACAT TTGGTTATGA AGAGCTAAAA GACAAGGAAA CAAACTACTT TTTTGTTGAA 3480 CCTTGGTTTG AGATACATAA TGAAGAGCTT GACTATGCTG TCCTGAAACT GAAGGAAAAT 3540 GGACAACAAG TACCTATGGA ACTATATAAT GGAATTACTC CTGTGCCACT TAGTGGGTTG 3600 ATACATATTA TTGGCCATCC ATATGGAGAA AAAAAGCAGA TTGATGCTTG TGCTGTGATC 3660 CCTCAGGGTC AGCGAGCAAA GAAATGTCAG GAACCTGTTC AGTCTAAAAA AGCAGAAAGT 3720 CCAGAGTATG TCCATATGTA TACTCAAAGA AGTTTCCAGA AAATAGTTCA CAACCCTGAT 3780 55 GTGATTACCT ATGACACTGA ATTITTCTTT GGGGCTTCCG GCTCCCCTGT GTTTGATTCA 3840 AAAGGTTCAT TGGTGGCCAT GCATGCTGCT GGCTTTGCTT ATACTTACCA AAATGAGACT 3900 CGTAGTATCA TTGAGTTTGG CTCTACCATG GAATCCATCC TCCTTGATAT TAAGCAAAGA 3960 60 CATAAACCAT GGTATGAAGA AGTATTTGTA AATCAGCAGG ATGTAGAAAT GATGAGTGAT 4020 GAGGACTTGT GAGAATTCAG TCTACTGGAT TTAAGGGAAT GGCTTATGGA GTTGFTATTT 4080 CGTAGGCATT GAAAATGGTT TTCTAAACTC CAAAATGGTC ATCTTATCAA TAATAATAAT 4140 ATTGACCATT TCCTATCTGC CAGGCATTTT TCTAAGCACA TGAAGAAATT AGTCCTAACA 4200 ACACTATGAG ATGGACTATA ACTTGCCCAA ATTTTTTTT TTTTTGAGAC TGAGTCTCAC 4260 65 70 GCAATGGGAA GAATAACAAG ATTATATAGT AATCAGTTTC ATGACACTAA AAGTCATATA 4620 GTCATAGGGT TTTTTCATCT TTCATATCTT TGCCTAAATT CATITGCTAC AGTGCAGGAA 4680 CCAAAACTTG TTCATCTCAT GATTCCCTAC ATCTGACATA AGGAAAGTAA GTGCTCAGAA 4740 AAATGTGCAG GTCAATAAGT TGCAAAAGTT GGGGCTGCAA TTAATGCTAA CATAAGAGCT 4800 AAATGCTTGA TTAGAAATGA TCTCAAAACC TTTTAGAATT TCCAAAATCT TCATATTACT 4860 75 GAAACTGTCG GAATATATGG GTCCTGAAAT TCAGAAGATG ATAGTCACTC TTCCCATATT 4920 TATAGGCTAT TAAGGCAAGG GATATCTTAA ACATCATATT ACTTTATTTA GATTTCTACT 4980 ACTCCAATTA TTAATGTTAT GTATTTCTCA TTGTTTTACT TCTTCATGGT ATTATGAAGA 5040 CTATATAGAT GATTCAACCA AGCCTGCAAA TCTCCCTCTT GTGGAATTCC ACTGGACCCA 5100 ATCTGTTTTC CATTTCCATT GCAATACTAC TAAAGCCATA CAATATCAAG CACCCTCCCT 5160

CTAGGTCCAG GGACTATCAC AGAAGAAGCA GGCATGTAAG ATTTTAAGGA CTGGTTTCGA 5220 GGGGTCGAGT GTAGGAAAAC AGCCTGTTGC ATTGTAAGAG TGATGTCACC TTGAAGAGCA 5280 GCTGGCATGA TGACTGCTGT TTGACTCCTG CATACCAAGA TATTCTGCAG CAATGTCTTT 5340 AAACAGTGCC GGTAGTACAG ATAACCCCTC ATAAAGATGC TTATCTAACC TCCCCAGTGT 5400 TCAGGTGTTT CACAAGAAAG TCTGAGCTAT GACTAGCTAC ACGTTTTCCC AAAAATGCTT 5460 GTTATATAAA GGGTACTTTT GGGAGGGTGA GTGCCGCCAT TTAGTGGCTG CTAGAAACAT 5520 TGCTTCTGTT TGTAAGTTCC TATTAAATGT TCTTTCTGAG AAAAAAAAA

SEQ ID NO:259 PBM4 Protein sequence: PBM4 Protein sequence: BAB67788

5

MDTVMKQTHA DTPVDHCLSG IRKCSSTFKL KSEVNKHETA LEMQNPNLNN KECCFTFTLN 60
GNSRKLDRSV FTAYGKPSES IYSALSANDY FSERIKNQFN KNIIVYEEKT IDGHINLGMP 120
LKCLPSDSHF KITFQQRKSES IYSALSANDY FSERIKNQFN KNIIVYEEKT IDGHINLGMP 120
KGSKLCIYAL KGETIEGALC KDGRIRSDIG EFEWKLKEGH KKIYGKQSMV DEVSGKVLEM 240
DISKKKALQQ KDHKKIKQN ESATDEINHQ SLIQSKKKVH KPKKDGETKD VEHSREQILP 300
PQDLSHYIKD KTRQTIPRIR NYYFCSLPRK YRQINSQVRR RPHLGRRYAI NLDVQKEAIN 360
LLKNYQTILNE AIMHQYPNFK EEAQWVRKYF REEQKRMNLS PAKQFNIYKK DFGKMTANSV 420
SVATCEQLTY YSKSVGFMQW DNNGNTGNAT CFVFNGGYIF TCRINVHLMV GKNTHPSLWP 480
DIISKCAKVT FTYTEFCPTP DNWFSIEPWL KVSNENLDYA ILKLKENGNA FPFGLWRQIS 540
PQPSTGLIYL IGHPEGQIKK IDGCTVIPLN ERLKKYPNDC QDGLVDLYDT TSNVYCMFTQ 600
RSFLSEVWNT HTLSYDTCFS DGSSGSPVFN ASGKLVALHT FGLFYQRGFN VHALIEFGYS 660
MDSILCDIKK TNESLYKSLN DEKLETYDEE KARPRPAYRR LGCFRFRSRF PILGTGETGR 720
EEAGKDRRGH GVSETGSCSR RQGGALWVSP AQPIGFRSSW SSGAFASSNT SGNCVERWIP 780
GRVLARRAVS KEQQNNCSTS LMRMESRGDP RATINTQAQR FHSPKKNPED QTMPQNRTTY 840
VTLKAVRKEI ETHQGQEMLV RGTEGIKEYI NLGMPLSCFP EGGQVVTIFS QSKSKQKEDN 900
HIFGRQDKAS TECVKFYIHA IGIGKCKRRI VKCGKLHKKG RKLCVYAFKG ETIKDALCKD 960
GRFLSFLEND DWKLIENDDT ILESTQPVDE LEGRYFQVEV EKRMVPSAAA SQNPESEKRN 1020
TCVLREQIVA QYPSLKRESE KIIENFKKKM KVKNGETLFE LHRTTIFGKVT KNSSSIKVVK 1080
LLVRLSDSVG YLFWDSATTG YATCFVFKGL FILTCRHVID SIVGDGIEPS KWATIIGQCV 1140
RVTFGYEELK DKETNYFFVE PWFEHNEEL DYAVLKLKEN GQQVPMELYN GITPVPLSGL 1200
HIIIGHPYGE KKQIDACAVI PQGQRAKKCQ ERVQSKKAES PEYVHMYTQR SFQKIVHNPD 1260
VITYYDTEFFF GASGSPVFDS KGSLVAMHAA GFAYTYQNET RSIIEFGSTM ESILLDIKQR 1320
HKPWYEEVFV NQQDVEMMSD EDL

#### SEQ ID NO:260 PBQ1 DNA sequence

Nucleic Acid Accession#: NM_015642
Coding sequence: 489-2489 (uni

Coding sequence: 489-2489 (underlined sequence corresponds to start and stop codon)

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		GAACTCTGAG					120
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	TGCAGCCGCT	CTCTGCTCCC	TGCCCCAATG	AACATCTGCA	CTAGGCCCAA	GCCTTGGAGT	240
50		AAGAGTGACA					300
50		GGCATCTGAG					360
	CGGGCCTTCC	CTGCCTGAAC	TTTGAAGCTG	TTTTGTCTCC	AGACCCAGCC	CTCATCCACT	420
	CAACACATTC	ACTGACAAAC	TCTCACGCTC	ACACCGGGTC	ATCTGATTGT	GACATCAGTT	480
	GCAAGGGGAT	GACCGAGCGC	ATTCACAGCA	TCAACCTTCA	CAACTTCAGC	AATTCCGTGC	540
ہے ہے	TCGAGACCCT	CAACGAGCAG	CGCAACCGTG	GCCACTTCTG	TGACGTAACG	GTGCGCATCC	600
<i>5</i> 5	ACGGGAGCAT	GCTGCGCGCA	CACCGCTGCG	TGCTGGCAGC	CGGCAGCCCC	TTCTTCCAGG	660
	ACAAACTGCT	GCTTGGCTAC	AGCGACATCG	AGATCCCGTC	GGTGGTGTCA	GTGCAGTCAG	720
		CATTGACTTC					780
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<i>(</i> 0	GCATCGTGTC	ACAGAACGTG	GGCGATGTGT	TCCCGGGGAT	CCAGGACTCG	GGCCAGGACA	900
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	ACCTGCAGAG	CCACCCACAG	CACAGCGTGG	ACAGGATCTA	CTCGGCACTC	TACGCGTGCT	1020
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		CGGCCTGCCC					1140
C =		CTCGCAGCAG					1200
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	AAGGTGAAAG	CTTCGACTCG	GGCGTCAGCT	CCTCCATAGG	CACCGAGCCT	GACTCGGTGG	1440
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	CCTCTCCGGA	GAGAAGCAAT	GAAGTGGAGA	TGGACAGCAC	TGTTATCACT	GTCAGCAACA	1620
•	GCTCCGACAA	GAGCGTCCTA	CAACAGCCTT	CGGTCAACAC	GTCCATCGGG	CAGCCATTGC	1680
	CAAGTACCCA	GCTCTACTTA	CGCCAGACAG	AAACCCTCAC	CAGCAACCTG	AGGATGCCTC	1740
~~	TGACCTTGAC	CAGCAACACG	CAGGTCATTG	GCACAGCTGG	CAACACCTAC	CTGCCAGCCC	1800
<i>7</i> 5	TCTTCACTAC	CCAGCCCGCG	GGCAGTGGCC	CCAAGCCTTT	CCTCTTCAGC	CTGCCACAGC	1860
	CCCTGGCAGG	CCAGCAGACC	CAGTTTGTGA	CAGTGTCCCA	GCCCGGTCTG	TCGACCTTTA	1920
	CTGCACAGCT	GCCAGCGCCA	CAGCCCCTGG	CCTCATCCGC	AGGCCACAGC	ACAGCCAGTG	1980
	GGCAAGGCGA	AAAAAAGCCT	TATGAGTGCA	CTCTCTGCAA	CAAGACTTTC	ACCGCCAAAC	2040
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80	GTTGGCGCTC	CTTCTCCTTA	AAGGATTACC	TTATCAAGCA	CATGGTGACA	CACACAGGAG	2160

	TGAGGGCATA	CCAGTGTAGT	ATCTGCAACA	AGCGCTTCAC	CCAGAAGAGC	TCCCTCAACG	2220
			GGAGAGAAGT				2280 2340
_			GAGCGACACG GGTGCCCGCG				2400
5	AGGGGACCAC	TTACGTCTGC	TCCGTCTGCC	CAGCAAAGTT	TGACCAAATC	GAGCAGTTCA	2460
•			GTGTCTGACG				2520 2580
			TTACTTTCTG				2640
10			CCTGCTGGT				2700
IO			CGGATGGTGG ACTGGCCTCT				2760 2820
	AAAAAAAAA	, microsmoc.					
15		SEQID	NO:261 PBQ1 Pr	otein sequence:			
	PBQ1 Protein sec	puence: NP_056	6457				
	MTERIHSINI	HNESNSVI ET	I NEORNRGHE	CDVTVRIHGS	MIRAHROVL	A AGSPFFQDKL	60
••	LLGYSDIEIP S	VVSVQSVQK	LIDFMYSGVL	RVSQSEALQI :	LTAASILQIK T	VIDECTRIV 120	)
20	SQNVGDVFPC	IQDSGQDTPI	R GTPESGTSGC	SSDTESGYLO	SHPQHSVDR	YSALYACSMQ	180
						LS TTPETTHCRK A EGTESEPKGE	
	SFDSGVSSSI (	TEPDSVEQQ	FGPGAARDSQ	AEPTQPEQAA	EAPAEGGPQT	NQLETGASSP	360
25						TSNLRMPLTL   QPGLSTFTAQ	
25	LPAPQPLASS	<b>AGHSTASGQC</b>	EKKPYECTLO	NKTFTAKQN	Y VKHMFVHT	GE KPHQCSICW	R 540
	SFSLKDYLIK	HMVTHTGVR	A YQCSICNKR	FTQKSSLNVH	M RLHRGEKS	YE CYICKKKFSI	I 600
	MRMHVSDG	, HSASNGIPPA	GIPPGARAG	PPGVVACIEG	1111103107	K FDQIEQFNDI	1 000
30			•				
	SEQ ID NO: 262	PROS DNA segue	nce				
	Nucleic Acid Acce	ssion#: Al6541	87	•			
35	Coding sequence	: 1-912 (	underlined sequen	ce corresponds to	start and stop cod	(na	
<i>JJ</i>	1	11	21	31	41	51	
	1	. !	· l	1		1	60
			CATATETTAC GTCGTTTCGA				120
40	CGGAAAACTC	CCTCACGATO	TAAAACGAAG	ATCAGGAGCA	GATTTGAAGA	ATTACAAAGT	180
			GTCAGAGACA ATTAAAGGAA				240 300
	AGCAGCAAAT	TAGAAAATGA	GTCCAAACTA	TTGTCATTAA	ACACTGATAA	AACTTTATGT	360
45			TCGAATTGAA				420 480
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			TGATACAGAC				660 720
50	TTTGCAGAAC	ATCGAGTACC	AAATGGAATG	AATAAGGGAG	AACATGCATI	AGTTCTGTTT	780
			ATATTTGCAG AAGAACCTCA				840 900
	TTATATAGAT		Monnecie	. MIIONCHOG	· imilico.		,,,,
55							
23	SEQ ID NO:263 F	BO6 Protein sea	ience:				
	Protein Accession						
	MERKEATCKI	NMVTKKKN	A FIRSRI YM	LE RRKTDTV	VES SVSGDHS	GTL RRSQSDRTI	EY 60
60	NOKLQEKMT	P QGECSVAET	L TPEEEHHMI	CR MMAKREK	IIK ELIQTEKD	YL NDLELCVRE	V 120
					DVEPAMQVIO	EVFLQIKOPL	180
	EDIYKIYCYH	HDEAHSILES	YEKEEELKEH	LSHCIQSLA			
<i></i>	•		•				
65	SEQ ID NO:264 F Nucleic Acid Acce						
	Coding sequence		725 (underlined sec	quence corresponi	is to start and stop	codon)	
					41		
70	1	11 	21 .	31 	41	51 }	
			GCCGCCGCC				60
			GCCCGCGCCA				120 180
75	GGGGACGCAG	CCCCCCCCC	CAGCGGGCCC	GGGAAAAGCC	GCGCGCGCG	CGCGCGCCTG	240
75			CCTCCCCGCG				300
	GCTCGCGCAC	CCCCCCGGAA	GGAGGTGGCA GGTAGACCGG	GAAGGGGAGG	CGGGCGGGCG	GAGAGGAGAG	360 420
	AGTGGCGCGC	AGTCCAGCGA	GGGCGGGGT	TGGCTATGTG	GGGGGTGGTG	CACCCCGCAG	480
80			CTGGGGGCGT GGGAGTGTGG				540
J	GCCICGGGCC	-arucaiaia	200101010G	veucate 10C	. 30300000		600

					TAGTGGGAGG		660
					TGCTACACAT		720
	CAGACACAGC	ACGGAGATGC	TGCACAACCT	GAACCAGCAG	CGCAAAAACG	GCGGGCGCTT	780
_	CTGCGACGTG	CTCTTGCGGG	TAGGCGACGA	GAGCTTCCCA	GCGCACCGCG	CCGTGCTGGC	840
5	CGCCTGCAGC	GAGTACTTTG	AGTCGGTGTT	CAGCGCCCAG	TTGGGCGACG	GCGGAGCTGC	900
	GGACGGGGGT	CCGGCTGATG	TAGGGGGCGC	GACGGCAGCA	CCAGGCGGCG	GGGCCGGGGG	960
	CAGCCGGGAG	CTGGAGATGC	ACACTATCAG	CTCCAAGGTA	TTTGGGGACA	TTCTGGACTT	1020
	CGCCTACACT	TCCCGCATCG	TGGTGCGCTT	GGAGAGCTTT	CCCGAACTCA	TGACGGCCGC	1080
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	CTCGGACTTG	GCCTTCCCTT	TGGACATGAC	CAACGGGGCA	GCCTTGGCAG	CCAACAGCAA	1260
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	GCTGGGCTAC	ATCGACCTTC	CTCCTCCGAG	GCTGGGTGAG	AATGGGCTAC	CCATCTCTGA	1680
	AGACCCCGAC	GGCCCCCGAA	AGAGGAGCCG	GACCAGGAAG	CAGGTGGCTT	GTGAGATCTG	1740
20	CGGCAAGATC	TTCCGTGATG	TGTATCATCT	TAACCGGCAC	AAGCTGTCCC	ACTCTGGGGA	1800
	GAAGCCCTAC	TCCTGCCCTG	TGTGTGGGTT	GCGGTTCAAG	AGAAAAGACC	GCATGTCCTA	1860
	CCATGTGCGG	TCCCATGATG	GGTCCGTGGG	CAAGCCTTAC	ATCTGCCAGA	GCTGTGGGAA	1920
	AGGCTTCTCC	AGGCCTGATC	ACTTGAACGG	ACATATCAAG	CAGGTGCACA	CTTCTGAGCG	1980
	GCCTCACAAG	TGTCAGACCT	GCAATGCTTC	TTTTGCCACC	CGAGACCGTC	TGCGCTCCCA	2040
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	ATACATGGCA	GACCACCTGA	AGAAGCACAG	CGAGGGGCCC	AGCAACTTCT	GCAGTATCTG	2160
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••	CTGCGCCAGG	ACCTATGGCA	ACAAAGAAGG	CCAGAAATGC	TCACATCAGG	ATCCGATTGA	2340
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	GAACAAACAC	ATCCAGAAGG	TGCATGTCCG	GGCTCTCGGG	GCCCCCTGG	GGGACCTGGG	2580
0.5						AGTCCTTTGG	2640
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						CCACCACCCA	2880
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•						TGAGGACACC	3360
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						ATTGTGTACC	3720
					GATGIGAATT	GATCTGATCA	3780
55	GACTGTATTA	AAAATGTTAG	TACATTACTC	TA			
رر							
	050 ID NO.005	DD1/7 D					
		PBY7 Protein se					
	Protein Access	ion#: NP_1	14439				

MERVNDASCG PSGCYTYQVS RHSTEMLHNL NQQRKNGGRF CDVLLRVGDE SFPAHRAVLA 60
ACSEYFESVF SAQLGDGGAA DGGPADVGGA TAAPGGGAGG SRELEMHTIS SKVFGDILDF 120
AYTSRIVVRL ESPPELMTAA KFLLMRSVIE ICQEVIKQSN VQILVPPARA DIMLFRPPGT 180
SDLGFPLDMT NGAALAANSN GIAGSMQPEE EAARAAGAAI AGQASLPVLP GVDRLPMVAG 240
PLSPQLLTSP FPSVASSAPP LTGKRGRGRP RKANLLDSMF GSPGGLREAG ILPCGLCGKV 300
FTDANRLRQH EAQHGVTSLQ LGYIDLPPPR LGENGLPISE DPDGPPKRSR TRAQVACEIC 360
GKIFRDVYHL NRHKLSHSGE KPYSCPVCGL RFKRKDRMSY HVRSHDGSVG KPYICQSCGK 420
GFSRPDHLNG HIKQVHTSER PHKCQTCNAS FATRDRLRSH LACHEDKVPC QVCGKYLRAA 480
YMADHLKKHS EGPSNFCSIC NREGQKCSHQ DPIESSDSYG DLSDASDLKT PEKQSANGSF 540
SCDMAVPKNK MESDGEKKYP CPECGSFFRS KSYLNKHIQK VHVRALGGPL GDLGPALGSP 600
FSPQQNMSLL ESFGFQIVQS AFASSLVDPE VDQQPMGPEG K 60 65 70

# SEQ ID NO:266 PBY9 DNA sequence Nucleic Acid Accession#: NM_012429

75 Coding sequence: 174-1385 (underlined sequence corresponds to start and stop codon)

	1	11	21	31	41	51	
	1		1	1			
00	CCCTACTCCG	CCTCTCGGGA	TCCTTTAAGA	GGCGGGGCTT	GGCTGCCAGC	TCCGCGGCCC	60
80	GGGCAAAAGG	CTGGGACTTT	ACTCCGGGTG	GCGGCGAGGA	CGAGTCTGTG	CTCCATCAGC	120

			GCCCCAAAC				180
			CCCAGGCAGA				240
			CTGCCGAATC				300
_			CAGAAGTCGG				360
5			AACATCATTA				420
			GGCTATGACC				480
			GGTCTGCTGT				540
			CTGCTTCTGC				600
10			ACCATAATTT				660
10			GCCTATGGAG				720
			TTTGTTGTTA				780
			AGTGAGGACA				840
			AAACATATCA				900
	GCACCATGAC	TGACCCTGAT	<b>GGAAACCCCA</b>	AGTGCAAATC	CAAGATCAAC	TACGGGGGTG	960
15	ACATCCCCAG	GAAGTATTAT	GTGCGAGACC	AGGTGAAACA	GCAGTATGAA	CACAGCGTGC	1020
	AGATTTCCCG	TGGCTCCTCC	CACCAAGTGG	AGTATGAGAT	CCTCTTCCCT	GGCTGTGTCC	1080
	TCAGGTGGCA	GTTTATGTCA	GATGGAGCGG	ATGTTGGTTT	TGGGATTTTC	CTGAAGACCA	1140
	AGATGGGAGA	GAGGCAGCGG	GCAGGGGAGA	TGACAGAGGT	GCTGCCCAAC	CAGAGGTACA	1200
	ACTCCCACCT	GGTCCCTGAA	GATGGGACCC	TCACCTGCAG	TGATCCTGGC	ATCTATGTCC	1260
20			AGCTTCATTC				1320
			TCAGAAGAGA				1380
			CAGGCCTGGC				1440
			CACAACCCTG				1500
			GTTAGGCAGA				1560
25			CCCAGGAGCT				1620
			ACCTGTCCAG				1680
			GGGAAAAAA				1740
			CGGGGAGAAA				1800
			GTAGCTGGTT				1860
30			GGCTGGGGTA				1920
50			TTCCCACTCG				1980
			CACACGGCCT				2040
			CGGGTACCCA				2100
			CGCAGCTGCA				2160
35			CCAGTGCCCT				2220
33			GAGACAAAAA				2280
			GAGTGTCCCG				2340
	GAGAGGGIGT	TIGCCAGICT	CAGTCCCATC	TOG TGCCCGC	CAACCCGCII	CCTGACTGAC	2400
							2460
40			AGCCAGGCCT				2520
40			GACTAGGGC				
	CAGCCCTTAC	CCCAATCCCA	CGAGCCCCGC	CAACGAACCA	CAGGTGCTGG	GCTTTAGAGA	2580
	ACATGGGAAG	GCGGCCCCAG	ACCTGGCGGG	AACGCCTTTC	CCTCAGAGCC	AGGCCCCGGC	2640
			TGCGAAGCTG				2700
AE			CTGCACGGGC				2760
45	TGGGTTTACA	ACGCTGTTAG	GAAAATTAAC	CAATGAATAA	AGCAACGTTC	AGTGCGCA	
		<u>PBY9 Protein se</u>					
50	Protein Accessi	on #: NP_03	6561				
50							
	MSGRVGDLS	PRQKEALAKI	R ENVQDVLP.	AL PNPDDYFL	LR WLRARSFI	OLQ KSEAMLRI	CHV 60
	EFRKQKDIDN	I IISWQPPEVI (	QQYLSGGMCC	S YDLDGCPVW	Y DIIGPLDAK	G LLFSASKQDI	L 120
	LRTKMRECE	LLLQECAHQT	T KLGRKVETI	T IIYDCEGLGI	L KHLWKPAVI	EA YGEFLCMFE	Œ 180
	NYPETLKRLE	· VVKAPKLFP	/ AYNLIKPFLS	EDTRKKIMVI	L GANWKEVLI	LK HISPDQVPV	E 240
55	YGGTMTDPD	G NPKCKSKIN	Y GGDIPRKY?	YV RDQVKQQ`	YEH SVQISRGS	SSH QVEYEILFT	G 300
	CVLRWQFMS	D GADVGFGI	L KTKMGERQ	RA GEMTEVL	PNQ RYNSHL\	PED GTLTCSD	PGI 360
	YVLRFDNTY	S FIHAKKVNF	r vevllpdka	S EEKMKQLG.	AG TPK		
							•
60	SEQ ID NO:268	PBH8 DNA seque	108				
		ession#: XM_00					
	Coding sequence		40 (underlined sec	quence correspond	is to start and stop	codon)	
			• • • • • • • • • • • • • • • • • • • •	•			
	1	11	21	31	41	51	
65	ī	1		1		1.	
	GTGGGGACAG	CCGAGCCGCG	CCGGGCCCCT		GCCAAGGAGC	TGGGATCGCA	60
			GATTTGTTTT				120
			TCCATTTAGG				180
			CTTCTGACCA				240
70			TGCTCCAAGG				300
. •	AUCHA A AUCTO	TCTTGGCGAA	AAGGAACGCG	GGCCTGACCT	GCAGCGGATA	CAAGGTCATC	360
			GATCAGGCAG				420
	ACCAPCAGE A	THE TEST TOTAL	GGTGGCCGTG	GGCCAGTCGC	TGCCACCCAG	TGCCATCACC	480
			CATGTTCATG				540
75	GUGATCHUCC	CCACCCOCAC	CUIGITONIA	CCCTACCACCA	CCCTTGWCCT	GATCGAGAAG	
13	*ICCIGOVII	PACTCACA	CGGCTGCGAC	CHCHACCYCC	TACCAGGACCA	ACACCACCTO	660
							720
			CACCACCAAG				
			CGCCACCGTG				780
90			TGTACTCACG				840
80	CTGGAGCAGG	TGTCCACTGC	CAAGTCCCAG	GACTCCTGGA	GGACCGCCTT	GICTACCICA	900

5	CAAGAAACTA ( AACCCTTACC ( CTCGGAAACT ( CACTCAGAAA (	CCCCACAGCA GGAGAGCCAG GCAGTGACCT	ATACAGCTCG ! ICCCCCTGCA ! ICTGTACACG (	TTCCAAATGG A AGCGCTGCTG ( CCATCCTACA (	ACAAACTGGA I CTCCTCCAGA I GCCTGCCCTT (	ATGCGGCCAG 1 ACTGCAGCCC 1 CTCCTACCAT 1	960 020 080 140
	TACGGACACT  AAGTTCGGGC  CCAGCCAGCG	AGCCCCAAGG / GTGAATGCCA (	ATCCCCTTGT ( GTGGCATTAT (	BAGGTGGCAC ( BCCAACCCCC !	GCTTTTTCCT (	SAGCACACTG 1 CAGCTCGTCT 1	200 260 320
10	CCAGCTAAAA TACGAAGGCA CTCCTGTTTG	AGCAGATGTC (	CTCTGCGGAG I				380 440
15	SEQ ID NO:269 P Protein Accession						
15	GDAWGQPSR/	GPLDGVAKE	L GSHLLQTLD	G FVFVVASD	GK IMYISETAS	KMRAVFPEGL IV HLGLSQVELT I RNAGLTCSGY	120
20	KLIFLDSRVT H RGGWVWVQS STSQETRKLV	EVTGYEPQDL I Y ATVVHNSRS KPKNTKMKTK	EKTLYHHVH SS RPHCIVSVN K LRTNPYPPQ(	GCDVFHLRYA Y VLTEIEYKE YSSFQMDKL	NHLLLVKGQ EL QLSLEQVST E CGQLGNWR	MFMFRASLDL V TTKYYRLLSK A KSQDSWRTAI AS PPASAAAPPI	300 2 360 3 420
25	STLPASGECQ Y PSFPSCGHYR I	WHYANPLVPS EEPALGPAKA	SSSPAKNPPE ARQAARDGAI	PPANTARHSL R LALARAAPE	VPSYEAPAAA C CAPPTPEAP	SPCEVARFFL 4 VRRFGEDTAP G APAQLPFVLL G APLPHYLGAS	540 600
30	Nucleic Acid Ac	SEQ (D ) cession#:	NO:270 PBJ9 DNA AA760894	A sequence;			
35	CCAGCCATGT GGAAACTCCA GGTGATGGAT	GGAACTGTTT TAGACCTTGT CTCTGCAGTA	TTCAGGTGCT TCCACTGGAA AAGTGGAAGA	G GTTCCATGO AC TCGTTCCCA AG TTCTTCAT	GC TCTTCCTG AT CTACCCTC GG CCCCCAA	FTG TGAGGCCTO AG CCGAAAATA CA CTCTATCCA GGT TATATCCA TA AGTTAGGTA	A 120 G 180 TC 240
40	AACTGAAGAT TAAAAAACAC GAGATTGGAC GAAATGAGGC TITGGACTTG	GAGATCATA GAAAGAACC GGATGCAGC GATTCTCCT CCCATAGCTT	C TGGATTAGO CC ATAGAAAC CC ACCGGCCC F AGAACCTTT GTATACTCTT	GA TGGGATCT AC AAGGAAG AG GAATGCC. A GAGAGRAC T ACTTTGGAT	FAA ATCCAAT GAAG GTCATG AGC AGCCACC AT GGTCCTGT A CAATTTTAT	GAA AATGTCTT TGAA GATGGA( CCAG AAGCTGG TGA ACAGCTTG. CCAAACTTGG(	CA 360 GGCA 420 AAG 480 AT 540 C 600
45	GATTCAAAGC AAAATATGAA TACACATGAA YGTGATCATY	AAGAAAATG GTGAACATT AACCCCCAA TAGAGATGT	A TGGGAACA G TGGTAGCTT G GGGAATCC A CAGAAAAG	TA GGAGGAG T AAGATGTT CC ATATCACA GT GAATCTGT	ACC AAGAAA TA GTGTAGCT AGT GTAGTGT IGT TCTGTAT	CTC GATTCTTA( GCCT ATAAAAA IGC AGGCACCC GAT ATTTGACA ATT CTGCCTAA(	AGCA 720 TA 780 TT 840 GG 900
<b>5</b> 0	GAAAACTGTA ATCTTCTTAC	AGCTTCCCAT TTGGACATTT AACACTTCCC	TATCTGGAGC CATGTGTTTA GCAGAGCTA	A TITCACITI GGGATTGTY A ACCATTAC	I'A AATATTTG T TYTAAATTC AG ANTATGA	AA AGCTITGCT GA TAAATATGI TI TCCTAATICA AAT AAAGACCC AAAAT GA	T 1020 1080
55	SEQ ID NO:271 P Nucleic Acid Acces Coding sequence:	sion#: AA14957	79	ce corresponds to	start and stop cod	ion)	
	1	11	21	31	41 i	51	•
60	GGCATAAATG	GTATCAAAGA	TGCAAGGAAG	AAGAGCCTTA GTCACTGTAG AGATGCGGCT	GTGAAACTTG GTGTGATTGG	AAGTGGAGAT	60 120 180
				CCTCATGTGG GTTGCTATAC			240 . 300
65	CTGTGGGACC	TGAGACATCT	GCTTGTGGGT	AAAATCCTGA	TTGATGTGAG	CAATAACATG	360
				GAATATTTGG GCTTGGGCAC			420 480
	GCCAGCCGGC	AGGTTTATAT	ATGCAGCAAC	AATATTCAAG	CGCGACAACA	GGTTATTGAA	540
70	ATTGAAAATT	TACCCCTACG	ACTOTTTACT	GACTTGGGAT CTCTGGAGAG	GGCCAGTGGT	GGTAGCTATA	600 660
				TTTGTCAGAG			720

AGCTTGGCCA CATTTTTTT CCTTTATTCC TTTGTCAGAG ATGTGATTCA TCCATATGCT AGAAACCAAC AGAGTGACTT TTACAAAATT CCTATAGAGA TTGTGAATAA AACCTTACCT

ATAGTTGCCA TTACTTTGCT CTCCCTAGTA TACCTCGCAG GTCTTCTGGC AGCTGCTTAT

ATACTTATT ACGCACCAA GTATAGGAGA TITCCACCIT GGITGGAAAC CIGGITACAG
TGTAGAAAAC AGCITGGATT ACTAGATITT TICTITCGCTA TGGTCACAGC
CTCTGCTTAC CGATGAGAAG GTCAGAGAGA TATTTGTTC TCAACATGGC TIATCAGCAG
GTTCATGCAA ATATTGAAAA CTCTTGGAAT GAGGAAGAAG TTTGGAGAAT TGAAATGTAT
ATCTCCTTTG GCATAATGAG CCTTGGCTTA CTTTCCCTCC TGGCAGTCAC TICTATCCCT

TCAGTGAGCA ATGCTTTAAA CTGGAGAGAA TTCAGTTTTA TTCAGTCTAC ACTTGGATAT 1200

	GAAGAGTAC		A TACACCACCI	A AACTTTGTT	TTGCTCTTG	AGCTTTTGAG TTTGCCCTCA	1260 1320
5	SEQ ID NO:272   Protein Accession	PBQ4 Protein sequ n #: none	uence;				
10	1	11	21	31	41	51	
10	RNPKFASEFF RINQYPESNA	KSLSETCLPN PHVVDVTHHE EYLASLFPDS DLGSLSSARE	DALTKTNIIF LIVKGFNVVS	VAIHREHYTS AWALQLGPKD	LWDLRHLLVG ASRQVYICSN	KILIDVSNNM NIQARQQVIE	60 120 180 240
15	CRKQLGLLSF ISFGIMSLGL	PIEIVNKTLP FFAMVHVAYS LSLLAVTSIP NFVLALVLPS	LCLPMRRSER SVSNALNWRE	YLFLNMAYQQ FSFIQSTLGY	VHANIENSWN	EEEVWRIEMY	300 360 420
20				SEC	ID NO:273 PBQ	DNA SEQUENCE	
	Nucleic Acid According sequence		1973 45 (underlined sec	wence correspond	s to start and stop	codon)	
25	1	11	21	31	41	51	
25	OCCOCCOUNT	CTACTCCGCC	COCCCCC		CCCCCCTCCTCCTCCTCCCTCCCTCCCTCCCTCCCTCC	TO CA CONTROC	60
		GGAGGCTGAG					120
		CGCGGCGTCG					180
20		CCTGCAGAAG					240
30		GCTTTTGCAG					300
		GAATTATGAC					360
		AGTGAATGGT TCCAATGACA					420 480
		CAGCAGTTCC					540
35		GACCTCTAGC					600
		TTTGAACTCC					660
		ACTGGCAGAG					720
		ACCTTCCAAA					780
40		TTCTCCATCT					.840 900
70		TTCCCTGGAA					960
		TCACCCAGAC					1020
		GAATTTGTCT					1080
4 5		TAATTCATCA					1140
45		GAGCAGTGAT					1200
		ACCAGCATTT					1260
		CCACTTCTGG TAACACACTT					1320 1380
		GCTGGATGGA					1440
50		GCACTTGTGG					1500
		GAAGTGAGCA					1560
	TTTGCCATTC	CCCATTGAAA	ACATCTTTTT	AGGATTCTCT	TTGAATAGGA	CTCAAGTTGG	1620
		TAAAAATGCC					1680
55		TTTCCTTCCT					1740
<b>J</b> J		TTTTGGTGGG					1800
		ATTTGTGAAA					1860 1920
	AAAAAAAA		Igrammount	70000000000		**************************************	1,20
60					•		
	SEQ ID NO:274 F Protein Accession	PBQ5 Protein sequ NP_001				-	
	MDSAITLWON	LLOLLOKPO	N KHMICWTS	ID GOFKLLOA	EE VARLWGII	KN KPNMNYD	KLS 60
65	RALRYYYVKI ENGGKDKPPC	N IIKKVNGQK	F VYKFVSYPE D YIHSGLYSSI	I LNMDPMTV F TLNSLNSSN\	GR IEGDCESL1 / KLFKLIKTEN	VF SEVSSSSKDV I PAEKLAEKKS	120
						PENLSLEP 300	•
<b>~</b> ^						ASLTPAFFSQ 3	
70	TPULTPSPL LE PGPFSPDLQK		VAPLSPAR LQ	GANTLFQF PS	VLNSHGPF TL	SGLDGPST 420	) ·
				4			
75	Nucleic Acid Acce Coding sequence		921 60 (underlined seq			DNA SEQUENCE codon)	
	_				44		
	1	11	21	31	41	51	
	1	1	[	1		1	
					41	6	
					• 4.	~	

			TTGACCGAGA				60
•	AGATGGAACT	TTAGACCAAA	AATTATTGGA	AGATTTACAA	AAGAAAAAA	ATGACCTTCG	120
	CTATATATA	ATTOCACOATT	TCAGAGAAAA	CONSOCRATION	ጥልጥርር እ አጥርር	AAAACCAATT	180
_			ATCAGGTAAC				240
5	AACCACTCAA	GTTACTCAGT	TCATTTTGGA	TAACTACATT	GAAAGAGGAA	AAGGATCTGC	300
	<b>ጥጥርር እር</b> እ አጥ እ	CHARGES	AGCCAAGAAG	AATTACTICCC	ATTENCACTORS	CCCANAGACT	360
			CTTGTGGCAG				420
	CCAGAGTCGG	TTGCCAAGGA	AACAGGGTTC	TATCTTATAC	TGTACAACAG	GAATCATCCT	480
	TO ACTICACOTO	CAGTCAGACC	CGTATTTGTC	CACTICTEACT	CATATICGTAC	THEATERANT	540
10							
10	CCATGAAAGA	AATCTGCAGT	CAGATGTTTT	AATGACTGTT	GTTAAAGACC	TICICAATIT	600
	TCGATCTGAC	TTGAAAGTAA	TATTGATGAG	TGCAACATTG	AATGCAGAAA	AGTTTTCAGA	660
	ATTATIVITY CO.	AACTETECAA	TGATACATAT	PCC-dACC-databath	ACCUMUCCCC	מידע מיבי מיבי מידע	720
			AAAAAATAAG				780
	CCAGTTTAAG	AGGGGTTTCA	TGCAAGGGCA	TGTAAATAGA	CAAGAAAAAG	AAGAAAAAGA	840
15	አርር አልጥልጥልጥ	DADCARCETT	<b>GGCCAGATTA</b>	TOTALCOCAL	CTCCCAACAA	CCTPATTACTCC	900
			AAATGATGGA				
							960
	TGCCCTCATC	CGATACATTG	TTTTGGAAGA	AGAGGATGGT	GCGATACTGG	TCTTTCTGCC	1020
	AGGCTGGGAC	AATATCAGCA	CTTTACATGA	TCTCTTGATG	TCACAAGTAA	TYPEAARTY	1080
			CTTTACATTC				
20							1140
20	GTTTAAAAGA	ACCCCTCCTG	GTGTTCGGAA	AATAGTAATT	GCTACCAACA	TTGCGGAGAC	1200
	TAGCATTACC	ATAGATGATG	TCGTTTATGT	GATAGATGGA	GGAAAAATAA	AAGAGACGCA	1260
			TCAGTACAAT				1320
	CAAACAGAGA	AAAGGTCGAG	CTGGAAGAGT	TCAACCTGGT	CATTGCTATC	ATCTGTATAA	1380
	TGGTCTTAGA	GCAAGTCTTC	TAGATGACTA	TCAACTGCCA	GAAATTTTGA	GAACTCCTTT	1440
25			TAAAGATTTT				1500
	TAGATTAATG	GACCCACCAT	CAAATGAGGC	AGTGTTACTC	TCCATAAGAC	ACCTGATGGA	1560
	GCTGAACGCT	TTGGATAAAC	AAGAAGAATT	GACACCTCTT	GGAGTCCACT	TGGCACGATT	1620
	ACCCCCTTCAC	CCACATIATIC	GAAAAATGAT	declaration (CV)	CCACTCTTCT	CCTCCTTACA	1680
20			CTAGTCTCAG				1740
30	AAAAGAAAAG	ATTGCAGATG	CAAGAAGAAA	GGAATTGGCA	AAGGATACTA	GAAGTGATCA	1800
	CTTAACAGTT	GTGAATGCGT	TTGAGGGCTG	GGAAGAGGCT	AGGCGACGTG	GTTTCAGATA	1860
			AATATTTTCT				1920
	CATGAAAGGA	CAGTTTGCTG	AGCATCTTCT	TGGAGCTGGA	TTTGTAAGCA	GTAGAAATCC	1980
	TAAAGATCCA	GAATCTAATA	TAAATTCAGA	TAATGAGAAG	ATAATTAAAG	CTGTCATCTG	2040
35			TTGCTAAAAT				2100
55							
	GGTAAAAGTT	TACACAAAAA	CCGATGGCCT	GGTTGCTGTT	CATCCTAAAT	CTGTTAATGT	2160
	GGAGCAAACA	GACTTTCACT	ACAACTGGCT	TATCTATCAC	CTAAAGATGA	GAACAAGCAG	2220
			CAGAGGTTTC				2280
•							
40			ACGATCAGGA				2340
40	TCAGTCTCCA	GCAAGAATTG	CCCATCTTGT	TAAGGAATTA	AGAAAGGAAC	TAGATATTCT	2400
	TOTOCANGAG	AACATTCAAA	GTCCTCATCC	TOTACACTCC	ABTORCACTA	AATTCACACA	2460
			TTATAGACTT				2520
	GAACTTTCCG	CCACGATTCC	AGGATGGATA	TTACAGCTGA	CAGCTTTTCA	GGGGTGGTCT	2580
	GAAAAGCCAG	TTTGACAGCC	ATTCTTCATC	ΔΑΑΤΤΤΥΣΥΓΤΑ	<b>ПЛАЦИССИСС</b>	ATGCCAAACC	2640
45			ATGTGTAAGG				
73							2700
	ATGTGCATGA	CITGATGTTA	TATGTAGAGA	TATATATATA	TATATATATA	CCATAAAAGC	2760
	AATATGTTCT	CTGATCATAT	ACTCTGCTGT	GGTCATGCCC	ACTCTTTGGG	AGTATATTCC	2820
			GTACCACTTG				2880
50			TTGATGATAC				2940
50	ACAAGTGTCA	ATTAAGAATT	TGAACACAAC	CACATTTTTT	AAAATGAAAC	TTCTATCGGA	3000
	ACTEAAATTAA	THEOTOPICOTAL	TAAAGTCCAG	מ מדים מידידים מידי	AATCTACAAT	CTTABATCTC	
	1101101111111111		1101010010	***********		01111111	
		=					
	SEQ ID NO:276 PI	3 <b>Y3 Protein seq</b> ue	nce:				
	Protein Accession	#: BAA960	12				
55	I Iddil Macadion	. 0/4/000	<b>'-</b>				
JJ							
	IRNRSYIDRD S	EYLLQENEP D	<b>IGTLDQKLLE</b> I	DLQKKKNDLR	YIEMQHFREK	( LPSYGMQKEI	L 60
	TYOHOU INV	VISGETGCGK	TTOVTOFII D I	NYTERGKGSA	CRIVCTOPER	ISAISVAERV 1	120
						VSHIVLDEI 1	
	HERNLQSDVL	MTVVKDLLNI	FRSDLKVILMS	S ATLNAEKFSI	E YFGNCPMIH	I PGFTFPVVEY	240
60	LLEDVIEKIR Y	VPEOKEHRS (	DEKRGEMOGH	VNROEKEEKE	ATYKERWPD	Y VRELRRRYS	A 300
						LLMSQVMFKS	
	DKFLIPLHS LI	MPTVNQTQV E	KRTPPGVRK I	VIATNIAET SI	TIDDVVYV ID	GGKIKETH 42	X)
	FDTONNISTM :	SAEWVSKANA	KORKGRAGR	V OPGHCYHT.	YN GLRASLLI	DY QLPEILRT	PL 480
						PLGVHLARL	
65							
65	PVEPHIGKMI I	FGALFCCLD F	VLTIAASLS FI	KDPFVIPLG KE	EKIADARRK EI	LAKDTRSDH (	500
	LTVVNAFEGW	EEARRRGFRY	EKDYCWEVE	L SSNTLOMI I	IN MKGOFAFI	HLL GAGFVSSF	NP 660
						, VAVHPKSVN	
	EQTOFHYNWL	. IYHLKMRTSS	IYLYDCTEVS	PYCLLFFGGD	ISIQKDNDQE	TIAVDEWIVE	780
						KTQEKATPR 8	
70	NFPPRFODGY		/·				- / <del>-</del>
, 0	וינדגגעטטז	13					

SEQ ID NO:277 PBY6 DNA SEQUENCE

Nucleic Acid Accession#: AA464018
Coding sequence: 64-1669(underlined sequence corresponds to start and stop codon)

GATTTTATCC TGGAACATTA CAGTGAAGAT GGCTATTTAT ATGAAGATGA AATTGCAGAT 60 CTT<u>ATG</u>GATC TGAGACAAGC TTGTCGGACG CCTAGCCGGG ATGAGGCCGG GGTGGAACTG 120

CTGATGACAT ACTTCATCCA GCTGGGCTTT GTCGAGAGTC GATTCTTCCC GCCCACACGG 180
CAGATGGGAC TCCTGTTCAC CTGGTATGAC TCTCTCACCG GGGTTCCGGT CAGCCAGCAG 240
AACCTGCTGC TGGAGAAGGC CAGTGTCCTG TTCAACACTG GGGCCCTCTA CACCCAGATT 300 GGGACCCGGT GTGATCGGCA GACGCAGGCT GGGCTGGAGA GTGCCATAGA TGCCTTTCAG 360 5 AGAGCCGCAG GGGTTTTAAA TTACCTGAAA GACACATTTA CCCATACTCC AAGTTACGAC 420 ATGAGCCCTG CCATGCTCAG CGTGCTCGTC AAAATGATGC TTGCACAAGC CCAAGAAAGC 480 GTGTTTGAGA AAATCAGCCT TCCTGGGATC CGGAATGAAT TCTTCATGCT GGTGAAGGTG 540 GCTCAGGAGG CTGCTAAGGT GGGAGAGGTC TACCAACAGC TACACGCAGC CATGAGCCAG 600 GCGCCGGTGA AAGAGAACAT CCCCTACTCC TGGGCCAGCT TAGCCTGCGT GAAGGCCCAC 660 10 CACTACGCGG CCCTGGCCCA CTACTTCACT GCCATCCTCC TCATCGACCA CCAGGTGAAG 720 CCAGGCACGG ATCTGGACCA CCAGGAGAAG TGCCTGTCCC AGCTCTACGA CCACATGCCA 780 GAGGGCTGA CACCCTTGGC CACACTGAAG AATGATCAGC AGCGCCGACA GCTGGGGAAG 840 TCCCACTTGC GCAGAGCCAT GGCTCATCAC GAGGAGTCGG TGCGGGAGGC CAGCCTCTGC 900 AAGAAGCTGC GGAGCATTGA GGTGCTACAG AAGGTGCTGT GTGCCGCACA GGAACGCTCC 960 15 CGGCTCACGT ACGCCCAGCA CCAGGAGGAG GATGACCTGC TGAACCTGAT CGACGCCCCC 1020 AGTGTTGTTG CTAAAACTGA GCAAGAGGGTT GACATTATAT TGCCCCAGTT CTCCAAGCTG 1080
ACAGTCACGG ACTTCTTCCA GAAGCTGGGC CCCTTATCTG TGTTTTTCGGC TAACAAGCGG 1140
TGGACGCCTC CTCGAAGCAT CCGCTTCACT GCAGAAGAAG GGGACTTGGG GTTCACCTTG 1200
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Nucleic Acid Accession#: AF107493

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2820

2880

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						(Q NKFSPLPLK)	
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80	AQDGLKKRLG	KLKATLEQAH	<b>QALEEEKKKT</b>	VDLLCSIFFC TRFDQQCGEL	<b>EVAQQLWQGQ</b>	VVQAKKFSNV AYCVAGGLHK	480 540

ESDTHAVQIA LMALKMMELS DEVMSPHGEP IKMRIGLHSG SVFAGVVGVK MPRYCLFGNN 660 VTLANKFESC SVPRKINVSP TTYRLLKDCP GFVFTPRSRE ELPPNFPSBI PGICHFLDAY. 660 QQGTNSKPCF QKKDVEDGNA NFLGKASGID

5

### SEQ ID NO:287 PFD2 DNA SEQUENCE

Nucleic Acid Accession#: NM_000720
Coding sequence: 119-6664 (underlined sequence corresponds to start and stop codon)

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	1	1	1		į.		
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IJ		ATGATGATGA					180
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	TATTCTACTG	AATGTCTTCT	TGGCCATCGC	TGTAGACAAT	TTGGCTGATG	CTGAAAGTCT	2460
		CAGAAAGAAG					2520
55		AATAAAAAGA					2580
33							
		ATTGATGACT					2640
		GTAGGGGAAG					2700
	CGGACCCCGT	CCTCGAAGGA	TCTCGGAGTT	GAACATGAAG	GAAAAAATTG	CCCCCATCCC	2760
	TGAAGGGAGC	GCTTTCTTCA	TTCTTAGCAA	GACCAACCCG	ATCCGCGTAG	GCTGCCACAA	2820
60		CACCACATCT					2880
		GCAGAGGACC					2940
		GCCTTCACAG					3000
	TGGAGCTTTC	CTCCACAAAG	GGGCCTTCTG	CAGGAACTAC	TTCAATTTGC	TGGATATGCT	3060
	GGTGGTTGGG	GTGTCTCTGG	TGTCATTTGG	GATTCAATCC	AGTGCCATCT	CCGTTGTGAA	3120
65	GATTCTGAGG	GTCTTAAGGG	TCCTGCGTCC	CCTCAGGGCC	ATCAACAGAG	CAAAAGGACT	3180
		GTCCAGTGCG					3240
		CTGCAGTTCA					3300
		ACGGATGAAG					3360
70		GGGGATGTTG					3420
70		GACAACGTCC					3480
		GCGTTGCTGT					3540
		CGCGTGGAGA					3600
	CTICATGATG	AACATCTTTG	TGGGCTTTGT	CATCGTTACA	TTTCAGGAAC	MAGGAGAAAA	3660
75	AGAGTATAAG	AACTGTGAGC	TGGACAAAAA	TCAGCGTCAG	TGTGTTGAAT	ACGCCTTGAA	3720
75	AGCACGTCCC	TTGCGGAGAT	ACATCCCCAA	AAACCCCTAC	CAGTACAAGT	TCTGGTACGT	3780
	GGTGAACTCT	TCGCCTTTCG	AATACATGAT	GTTTGTCCTC	ATCATGCTCA	ACACACTCTG	3840
	Chilecus	CAGCACTACG	PCC PCACCA A	CATCTOVANT	GATGCCATCC	ACATTOTCAN	3900
		ACCGGGGTGT					3960
00		TTTAGTGACG					4020
80	TATAGACGTG	GCCCTCAGCG	AAGCGGACCC	AACTGAAAGT	GAAAATGTCC	CTGTCCCAAC	4080

	TGCTACACCT	GGGAACTCTG	AAGAGAGCAA	TAGAATCTCC	ATCACCTTTT	TCCGTCTTTT	4140
						CATTGCTGTG	4200
			AGGCGCTCCC				4260
_						GAGATAACAA	4320
5	CCAGATCAAT	AGGAACAATA	ACTTCCAGAC	GTTTCCCCAG	GCGGTGCTGC	TGCTCTTCAG	4380
	GTGTGCAACA	GGTGAGGCCT	GGCAGGAGAT	CATGCTGGCC	TGTCTCCCAG	GGAAGCTCTG	4440
	TGACCCTGAG	TCAGATTACA	ACCCCGGGGA	GGAGTATACA	TGTGGGAGCA	ACTTTGCCAT	4500
	TGTCTATTTC	ATCAGTTTTT	ACATGCTCTG	TGCATTTCTG	ATCATCAATC	TGTTTGTGGC	4560
4.0	TGTCATCATG	GATAATTTCG	ACTATCTGAC	CCGGGACTGG	TCTATTTTGG	GGCCTCACCA	4620
10	TTTAGATGAA	TTCAAAAGAA	TATGGTCAGA	ATATGACCCT	GAGGCAAAGG	GAAGGATAAA	4680
	ACACCTTGAT	GTGGTCACTC	TGCTTCGACG	CATCCAGCCT	CCCCTGGGGT	TTGGGAAGTT	4740
	ATGTCCACAC	AGGGTAGCGT	GCAAGAGATT	AGTTGCCATG	AACATGCCTC	TCAACAGTGA	4800
						TTAAGATCAA	4860
1.5			AAGCTAATGA				4920
15						ATGATGAGGT	4980
						AATTCAAGAA	5040
						TTGCCCTACA	5100
						TATCGTGTGA	5160
20						TGTTCAAAAG	5220
20						GAGATTCCCT	5280
			ACCGTCCCCT				5340
						ATAACCATCA	5400
			AGCAAGTTCC				5460
25			ATGGAAAGCG				5520
23			CCCACAAGCA				5580
			AAACTTACAT				5640
			CAGAGATACA			ACTGCTTGGG	5700 5760
30			ACAGCAGATA			TTTGCTATGA	5820 5880
50			GACGCCTACT				5940
						CGTCGTCTCC	6000
			CCCTGCCTCT				6060
			AAGCCCAGAA				6120
35						CCCCCTGAT	6180
-			CCCTGGACCA				6240
			AGCCCGACAT				6300
			GGAACAAAAA				6360
			CCGAAGGCTT				6420
40						AGATGGAGAG	6480
						ATGTGGGCCC	6540
	CCTCTCACAC	CGGCAGGACT	ATGAGCTACA	GGACTTTGGT	CCTGGCTACA	GCGACGAAGA	6600 ·
	GCCAGACCCT	GGGAGGGATG	AGGAGGACCT	GGCGGATGAA	ATGATATGCA	TCACCACCTT	6660
4 ~						AGAGCCAGGG	6720
45			GAAAGTTTAG				6780
			TCATGCCTCA				6840
						AAACAGCAGG	6900
						ATTGTCCAGA	6960
50						ACCCAACTGA	7020
JU						CACCTCGTGT	7080
						CATTTTTAAA	7140
	CCCTTCCCC	CAAATACACT	GCGTCCTGGT	TCCTGTTTAG	CIGITCIGAA	ATA	
	CEO ID NO.200 I	PFD2 Protein sequ			_		
55		#: A38198					
33	FIOLEIII ACCESSION	IW. A30130	,				
				24	44	51	
	1	11	21	31	41	1	
	MMMMMKKM		AMEDIANA DOM	RLPLSGEGPT	SOPNSSKOTV	LSWQAAIDAA	60
60			SORKROOYAK				120
00			VALAIYIPFP				180
			DFVIVIVGLF				240
			LNSIIKAMVP				300
			GNGRQCTANG				360
65			WPWVYFVSLI				420
			WITQAEDIDP				480
			AAKAGPSGCR				540
			SSEHYNOPDW				600
			ETILVELEIM				660
70			IIIFSLLGMQ				720
	ILTGEDWNAV	MYDGIMAYGG	PSSSGMIVCI	YFIILFICGN	YILLNVFLAI	AVDNLADAES	780
			ESLENKKNNK				840
			AGPRPRRISE				900
			AALAAEDPIR				960
<i>7</i> 5			LVVGVSLVSF				1020
			VTTLLQFMFA				1080
			DFNFDNVLSA				1140
			FFMMNIFVGF				1200
00						KMFNDAMDIL	1260
80	NMVFTGVFTV	EMVLKVIAFK	PKGYFSDAWN	TFDSLIVIGS	IIDVALSEAD	PTESENVPVP	1320
					42	_	

	TATPGNSEES	NRISITFFRL	FRVMRLVKLL	SRGEGIRTLL	WTFIKSFQAL	PYVALLIAML	1380
			NQINRNNNFQ				1440
						TRDWSILGPH	1500
5			KHLDVVTLLR				1560
5			KTEGNLEQAN KRKEQGLVGK				1620 1680
						LHVQRPSIPP	1740
						RPSIGNLEHV	1800
			VKRTRYYETY				1860
10						LEDDDSPVCY	1920
	DSRRSPRRRL	LPPTPASHRR	SSFNFECLRR	QSSQEEVPSS	PIPPHRTALP	LHLMQQQIMA	1980
			WATPPATPPY				2040
						LGRYARDPKF	2100
15			SAASTLLNGN	VRPRANGDVG	PLSHRQDYEL	QDFGPGYSDE	2160
13	EPDPGKDEED	LADEMICITT	ь				
				SF	O ID NO-289 ORIA	DNA SEQUENCE	
	Nucleic Acid Acc	ession#: NM_00	2812		<b>4</b> 10 110.200 00.0	DIVICEGOENCE	
	Coding sequence		62 (underlined sec	quence correspond	is to start and stop	codon)	
20	• • •			•		·	
	1 (4)	11	21	31	41	51	
	ī	ī	Ī	1	1	1	•
	AACTCCCGCC	TCGGGACGCC	TCGGGGTCGG	GCTCCGGCTG	CGGCTGCTGC	TGCGGCGCCC	60
25			CCTGTGCCCG				120
			CCCGCCGCGA				180
			AGCGTCCTGC				240
			CCGGGCCCGG				300 360
30			CCGGGCCCCG				420
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			TCCTTCAACA				540
•			GCTGAGATCC				600
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			TATTCCTGCT				780 840
			AGCATTGCTG				900
			TGGCTCTTTG				960
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	TCCGGCCACG	CAATGCAGGG	ATCTACCGCT	GCATTGGCCA	GGGGCAGAGG	GGCCCACCCA	1080
			CACCTAGCAG				1140
			GAGGAGCGTG				1200
45			CACGCGGGAG TTGGCCAATA				1260 1320
73			GGTCAGCGGA				1380
			CCCCAAGACA				1440
			ACACCAAAAC				1500
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			GTGCTGGAAA				1680
			AAGGAGGCCA CGGGCAGATG				1740 1800
			GCCCGGGTGA				1860
55			GGCCAGATTC				1920
			CCAGAGCGTA				1980 .
			GACCCCAAGC				2040
	TCCTGGACCC	CACCAAGCTG	GGACCCAGGA	TGCACATCTT	CCAGAATGGC	TCCCTGGTGA	2100
60			GACTCAGGCC				2160
UŲ			CCCTCTATG			TCGGTGGGTG	2220 2280
						AAGCGCTGCA	
						TGCCTCAACG	2400
						GCCTTGACCA	
65	GCTTGGGCTC	CGGCCCGCG	GCCACCAACA	AACGCCACAG	CACAAGTGAT	AAGATGCACT	2520
						GGGGAGGTGT	2580
						CTTGTGAAGA	2640
						GAGATGTTTG	
70						GCTGAGCCCC	2760
70						AGGATTTCCA GTGGCCCTAT	2820 2880
						CATAAGGACT	2940
			GTCAGTGCCC				3000
			AGTGAGTACT				3060
<b>75</b>						GATGTCTGGG	3120
						GGTGGGCAGG	3180
						CAGCCCGAGG	3240
						CCCAAGGACC	3300
80			ATGGCCTGGG			AGCAAGCCG <u>T</u>	3360 3420
50	<u>avannanun</u>	CCCGC I CAGG		HUUNUUUN	CHICICINGA		3460

5 10	TTGCTGAGGT GGCTGACTTG CTCTTCCTCT TTCTCCCCTT AGGCTTGGGA AGGGTTAATG ACACAGCAAG CCCCACCTTTGACAC TGCAGCGTGG GCCATCCTTA	GGCAAGATCC CTGAGCAGGG GACCCAAACT ATCAGGGACA GACCGGGTCC TGAGCTGGGT AGTCTCTCC TCTCCTTTC TATATAAACC GGTGGGTGGG CCCCACACTT TTTACACTCG	CCTGGCCTTT GGGCGACTAG GTGTTGGGTGC AACTCTGCCA TTGTGGGAG CCACTGGTCC CCACCTAGG CTCATCCTAA GCCCTTTTTG CATGGAGGT TTATTGTTGT	CCTCCTCTTC GGCTTTGAGC CACAGGTAAC CTCATCTGCC TTCCTTAATA ACTTGGGGT CTTGTGCACA TATGCACCAC AGGGGTGGGC CGTTTTTTGT	CTCACCCTCA TGGGCAGTTT CCCAATTTCT TTCTCAAGTT CTAGACCAGA CTGACCCAGA GATGAACGAG GGGCGGCTTT CCTTGAGATG TTGTTTTGTT	TCCTTTGGA CCCTGCCAC GCCCTCCAC GCGGAGGGCT CTGGGCACAC ATTATAGAGG CCCACGTCTT TTTTCAGGAA TATATGTAAT AGGAGGGTGG	3480 3540 3600 3660 3720 3780 3840 3900 3960 4020 4080 4140
15	SEQ ID NO:290 Protein Accessio	OBI6 Protein segu n #: NP_0	ence: 02812				
20	1	11	21	31	41	51	
20	VHVYWLLDGA	PRRLPLLSVL PVQDTERRFA LKHPASEAEI	<b>QGSSLSFAAV</b>	DRLQDSGTFQ	CVARDDVTGE	EARSANASFN	60 120 180
		GPEHSGLYSC					240
25		QPPPSLQWLF					300
		IILEATLHLA					360
		QKGHELVLAN					420
		LDCLTQATPK					480
30		BAQARVQVLE NAGTLHFARV					540
<b>J</b>		LOCEAOGDPK					600 660
		NIKHTEAPLY					720
		KAKRLQKQPE					780
25		FPRSSLQPIT					840
35		GKLNHANVVR					900
		CTQVALGMEH					960
		RWMSPEAILE GCPSKLYRLM				ADDEATADD	1020
•	MOIGHTHI ALD	GCLOILLINGI	Q.C.C.A.D.D.I.A.D	KE DI DILINDA	DGDS1 VDSRE		
40							
				SEC	2 ID NO:291 AAB	DNA SEQUENCE	
	Nucleic Acid Acc		NM_002205				
	Nucleic Acid Acc Coding sequence			SEC ined sequences co			
45							
45							
45	Coding sequence	: 11 	1-3150 (underl	ined sequences co	orrespond to start a	and stop codons) 51	
45	Coding sequence  1   ATGGGGAGCC	11       GGACGCCAGA	1-3150 (underl	ined sequences co	orrespond to start a	and stop codons) 51   GGGCCCCCGG	60
	Coding sequence  1   ATGGGGAGCC CGCCGACCCC	11     GGACGCCAGA   CGCTSSTGCC	1-3150 (under	ined sequences co	41   AGCTGCGCTG	and stop codons)  51   GGGCCCCCGG CAGGGTCGGG	60 120
45 50	Coding sequence  1   ATGGGGAGCC CGCCGACCCC GGCTTCAACT	11     GGACGCCAGA   CGCTSSTGCC   TAGACGCGGA	1-3150 (under	ined sequences co	41 	ind stop codons)  51   GGGCCCCCGG CAGGGTCGGG CTCCTTCTTC	60 120 180
	1   ATGGGGAGCC CGCCGACCCC GGCTTCAACT GGATTCTCAG	11     GGACGCCAGA CGCTSSTGCC TAGACGCGGA TGGAGTTTTA	1-3150 (under	31   CACGCCGTGC CTGCTSSTGC GTACTCTCGG ACAGACGGGG	41 AGCTGCGCTG GGCCCCCGGG TCAGTGTGCT	ind stop codons)  51   GGGCCCCCGG CAGGGTCGGG CTCCTTCTTC GGTGGGAGCA	60 120 180 240
	1   ATGGGGAGCC GGCTTCAACT GGATTCTCAG CCCAAGGCTA	11     GGACGCCAGA   CGCTSSTGCC   TAGACGCGGA	1-3150 (under	31   CACGCCGTGC CTGCTSCTGC ACAGACGCGG	41   AGCTGCGCTG GGCCCCCGGG TCAGTGTGCTA	sind stop codons)  51   GGGCCCCCGG CAGGGTCGGG CTCCTTCTC GGTGGGAGCA CCTCTGTCCT	60 120 180
50	1   ATGGGAGCC GGCTACACT GGATTCTCAG CCCAAGGCTA TGGGGTGCCA CTGGAGTCCT CTGGAGTCCT	11   GGACGCCAGA CGCTSSTGCC TAGACGCGGA TAGACTTTTA ATACCACCA GCCCACACA CACTGTCCAG	1-3150 (under	31   CACGCCGTGC CTGCTSSTGC GTACTCTCGG ACAGACGGGG ACTGCAGGGTG ATTGAATTTG GAGGAGCCTG	41   AGCTGCGCTG GGCCCCCGGG TCAGTGTGCT GTGCTGTCTGTTGTTAACAGCAAAAGG TGGAGTACAA	sind stop codons)  51   GGGCCCCCGG CAGGGTCGGG CTCCTTCTTC GGTGGGAGCA CCTCTTGTCCT CTCTCGGCTC GTCCTTGCAG	60 120 180 240 300 360 420
	1   ATGGGAGCC GGCTACACT GGATTCTCAG CCCAAGGCTA TGGGGTGCCA TGGGTTCCAT TGGTTCGGGG	11   GGACGCCAGA CGCTSSTGCC TAGACGCGGA TGGAGTTTTA ATACCAGCCA GCCCACACA GCCCCACAC CACTGTCCAG CAACAGTTCG	1-3150 (under	31   CACGCCGTGC CTGCTSSTGC GTACTCTCGG ACAGACGGGG ACTGCAGGGTG ATTGAATTTG GAGGAGCCTG TCCTCCATCT	41   AGCTGCGCTG GGCCCCCGGG TCAGTGTGCT GTGCTGTCTA ACAGCAAAGG TCGAGTACAA TGGAGTACAA	sind stop codons)  51   GGGCCCCCGG CAGGGTCGGG CTCCTTCTTC GGTGGGAGCA CCTCTGTCCT CTCTCGGCTC GTCCTTGCAG TCCACTGTAC	60 120 180 240 300 360 420 480
50	1 ATGGGGAGCC CCCCAACCCC GGCTTCAACT GCATTCTCAG CCCAAGGCTA TGGGGTGCCA TGGGGTCCT TGGTTCCGGGG AGCTGCCCA	11   GGACGCCAGA CGCTSSTGC TAGACGCGGA TGGAGTTTTA ATACCAGCCA GCCCCACCA CACTGTCCAG CAACAGTTCG CAGAGAAGGA	1-3150 (under	31   CACGCCGTGC CTGCTSSTGC GTACTCTCGG ACAGACGGGG ACTGCAGGGTG ATTGAATTTG GAGGAGCCTG TCCTCCATCT GACCCCGTGG	41   AGCTGCGCTG GGCCCCCGGG TCAGTGTGCTA ACAGCAAAGG TCGGAGTACAA TGGCATGCGC GCACCTGCTA	sind stop codons)  51   GGGCCCCGG CAGGGTCGGG CTCCTTCTTC GGTGGAGCA CCTCTGCGCTC CTCTCGGCTC GTCCTTCTGCAC CCCTCTCACAC	60 120 180 240 300 360 420 480 540
50	Coding sequence  1	11   GGACGCCAGA CGCTSSTGCC TAGACGCGGA TAGACGCA ATACCAGCA GCCCACACA CACTGTCCAG CACAGATTCG CAGAGAAGGA CCCGAATTCT	1-3150 (under 21 1 GTCCCCTCTC GCGCCGGCA CCGGCCGGGA GCCAGGAGTG GTGCACCCCC CTCAGAGGGA AGCCCATGGC GCCACTGAGC GCACTGAGC	ined sequences of  31     CACGCCGTGC CTGCTSSTGC GTACTCTCGG ACAGACGGGG CTGCAGGGTG ATTGAATTTG GAGGAGCCTG GACCCGTGG CCCTGCCCCTT	41   AGCTGCGCTG GGCCCCCGG TCAGTGTGCT ACAGCARAGG TGGAGTACAA TGGAGTACAA TGGAGTACAA TGGAGTACAC TGGACTGCTC CAGATTTCAG	sind stop codons)  51   GGGCCCCCGG CAGGGTCGGG CTCCTTCTTC GGTGGGAGCA CCTCTCGCCTC GTCCTTCCAG TCCACTCTACA CCTCTCCACA CTGGCCAGCA	60 120 180 240 300 420 480 540 600
50 55	1   ATGGGAGCC GGCTTCAACT GGATTCTCAG CCCAAGGCTA CTGGAGTCCT TGGTTCGGG AGCTGCGGG AGCTGCGCGT GGATACTTCA	11   GGACGCCAGA CGCTSSTGCC TAGACGCGGA TGGAGTTTTA ATACCAGCA GCCCACACA CACTGTCCAG CACAGTTCG CAGAGAAGGA CCCGAATTCT ACTGCCAAGG	1-3150 (under 21   GTCCCCTCTC GCCCCAGCA CCGGCCGGGA GCCAGCAGTG GTGCACCCC CTCAGAGGGA AGCCCATGGC GCACTGAGC GGAGTATGCA AGGCTTCAGT	ined sequences of  31   CACGCCGTGC CTGCTSSTGC GTACTCTCGG ACAGACGGGG CTGCAGGGTG ATTGAATTTG GAGGAGCCTG GACCCCGTGG CCCTGCCGCTTG	41   AGCTGCGCTG GGCCCCCGGG TCAGTGTGCT ACAGCAAAGG TGGATACAA TGGCATGCGC GCACCTGCTA CAGATTTCAG CCAAGACTGG	and stop codons)  51   GGGCCCCCGG CAGGGTCGGG CTCCTTCTTC GGTGGGAGCA CTCTTGCCT GTCCTTCCAG TCCACTGTAC CCTCTCCACA CCTGTCCACA CCTGGCAGCA CCGTGTGGTT	60 120 180 240 300 360 420 480 540 600 660
50	1   ATGGGGAGCC GGCTGACCCC GGATTCTCAG CCCAAGGCTA TGGGGTGCCT TGGTTCGGG AGCTGGCGCA AGCTGACTCT TGTTCGGG TTAGGTGAC	11   GGACGCCAGA CGCTSSTGCC TAGACGCGGA TAGACGCA ATACCAGCA GCCCACACA CACTGTCCAG CACAGATTCG CAGAGAAGGA CCCGAATTCT	1-3150 (under  21    GTCCCCTCTC GCTGCTGTTG GCCCCAGCA CCGGCCGGGA GCCAGGAGTG CTCAGAGGGA AGCCCATGACG GCACTGACG GGAGTATGAC AGGCTTCAGT TTTCTGGCAA	31   CACGCCGTGC CTGCTSSTGC GTACTCTCGG ACAGACGGGG ACTGCAGGGTG ATTGAATTTG AGAGACCTG TCCTCCATCT GACCCCGTGG CCCTGCCGCT CCCTGCCGCT GCCGAGTTCA GGCCAGATCC	41   AGCTGCGCTG GGCCCCCGGG TCAGTGTGCT ACAGCAAAGG TGGAGTACAA TGGCATGCAC GCACCTGCTA CAGATTTCAG CCAAGACTGG TGTCTGCCAC	sind stop codons)  51   GGGCCCCCGG CAGGGTCGGG CTCCTTCTTC GGTGGGAGCA CCTCTTGCAG TCCACTGTAC CCTCTCCACA CTGGCAGCA CCGTGTGGTT TCAGGAGCAG	60 120 180 240 300 360 420 480 540 600
50 55	Coding sequence  1	11   GGACGCCAGA CGCTSSTGCC TAGACGCGGA TAGACGCACA CACTGTCCAG CACAGATTCG CAGAGAAGTA CCGAATTCT ACTGCCAAGG CAGAGAACTA CTTATTACCC GTTCCATCTA	1-3150 (under  21    GTCCCCTCTC GCCCCGCAGCA CCGGCCGGGA GCCAGGAGTG GTGCACCCCC CTCAGAGGGA AGCCCATGGC GCACTGAGC GCACTGAGC GGAGTATGCA AGGCTTCAGT TTTCTGGCAA CGAGTACCTG TGATGACAGC	ined sequences or  31     CACGCCGTGC CACGCCGTGC CTGCTSSTGC GTACTCTCGG ACAGACGGGG ATTGAATTTG GAGGAGCCTG GACCCCGTTGACCTCGACCCGTGGCCGGAGTTCA GCCAGATCA ATCAACCTGG TACCTAGGAT	41  AGCTGCGCTG GGCCCCCGGG TCAGTGTGCT TCAGGCARAGG TGGAGTACAA TGGCATGCGC GCACCTGCTA CAGATTTCAG CCAAGACTGG TGTGCTGCTA CAGATTTCAG CCAAGACTGG TGTCTGCCAC TTCAGGGCA ACTCTGTGCC	sind stop codons)  51   GGGCCCCCGG CAGGGTCGGG CTCCTTCTTC GCTGGCTC GTCCTTGCAG TCCACTGCAG CCTCTCCACA CTGGCCAGCA CCGGCTGGCTT TCAGGAGCAG TCAGGAGCAG TCAGGAGCAG TCAGGAGCAG TCAGGAGCAG TCAGGAGCAG TCTTGGTGAA	60 120 180 240 300 360 420 480 540 660 720 780 840
50 55	1   ATGGGAGCC GGCTGACCCC GGCTTCAACT GGATTCTCAG CCCAAGGCTA CTGGAGTCCT TGGTTCGGG AGCTGCCGCA GGATACTTCA GGACAGGGTT TTAGGTGGAC ATTGCAGAACT TTAGGTGGAC ATTGCAGACT TTCAGTGGTCT	11   GGACGCCAGA CGCTSSTGCC TAGACGCGGA TGGAGTTTTA ATACCAGCCA CACTGTCCAG CAACAGTTCG CAGAGAAGGA ACCGCAATTCT ACTGCCAAGG CAGGAAGCTA CTTATTACCA GTTCCACG GTTCCACTA	1-3150 (under  21  GTCCCCTCTC GCTGCTGTTG GCCCCAGGA CCGGCCGGGA GCCAGGAGTA CTCAGAGGGA AGCCCATGAC GCACTGACC GCACTGACC TTTCTGGCAA AGGCTTCAGT TTTCTGGCAA TGATGACAGC AGACTTTCTT	ined sequences of  31   CACGCCGTGC CTGCTSSTGC GTACTCTCGG ACAGACGGGG CTGCAGGGTG TCCTCCATCT GACCCCGTGG CCCTGCCGCT GCCGAGTTCA GGCCAGATCC ATCAACCTGG TACCTAGGAT GCTGGTGTGC	41   AGCTGCGCTG GGCCCCCGGG TCAGTGTGCT ACAGCTACAA TGGCATACAA TGGCATACAA TGGCATTCAG CCAAGACTGG TGTCTGCCAC TTCAGGGGCA ACTCTGTGCAC TCTCTGCGCAC CCAAAGGGAA	and stop codons)  51    GGGCCCCCGG CAGGGTCGGG CTCCTTCTTC GGTGGGAGCA CCTCTTCCAG TCCACTGTAC CCTCTCCAG TCCACTGCAG CCTGCCAGA CCGGGAGCA CCGTGTGGTT TCAGGAGCAG CCGTGTGGTT TCAGGAGCAG CCTCACACA CCTCACACA CCTCACACA CCTCACACAC CCTCACACAC CCTCACACAC CCTCACACAC CCTCACACAC CCTCACACAC CCTCACACAC CCTCACACAC CCTCACACTTAC	60 120 180 240 300 420 480 540 600 660 720 780 840 900
50 55	1 ATGGGGAGCC GGCTATGTCAG GGATTCTCAG CCCAAGGCTA TGGGGTGCCT TGGTTCGGG AGCTGGCGCA AGCTGGCGCA AGCTGGCGCA AGCTGGCGCA ATTGGGTGAC ATTGGTGGAC ATTGGTGGAC ATTGGTGGAC ATTGCAGATT TTAGGTGGAC ATTGCAGATT GGCCAGGCCTA	11   GGACGCCAGA CGCTSSTGCC TAGACGCGA TGGAGTTTTA ATACCAGCCA CCCCACACA CCACTGTCCAG CAGAGAAGGA CCGGAATTCT CAGGGAAGCTA CTTATTACCC GTTCCATCTA ATGACACAGA CCATCCTTAA	1-3150 (under  21    GTCCCCTCTC GCTGCTGTTG GCCCCAGGA GCCAGGAGTG GTGACCCC GCACTGAGG AGCCCATGAG AGCCTTCAGT TTTCTGGCAA CGAGTACCTG TGATGACCAG AGCTTCAGT TGATGACAGC TGATGACAGC TGATGACAGC TGATGACAGC TGATGACAGC TGATGACAGC TGATGACAGC	31   CACGCCGTGC CTGCTSSTGC GTACTCTCGG ACAGACGGGG ACTGCAGGGTG CTGCAGCTG GAGGAGCCTG GAGCCCTGCAGCTG GCCCTGCCGCT GCCGAGTTCA GCCAGATCC ATCAACCTGG TACCTAGGGT GCTGGTGTGC ATCAACCTAGGA TACCTAGGAT ACCTAGGAT ACCTAGGATCC ATTCGATCCC	41   AGCTGCGCTG GGCCCCCGGG TCAGTGTGCT ACAGCTACAA ACAGCAAAGG TCGAGTACAA TGGCATGCTA ACAGCATGCTA CAGATTTCAG CCAAGACTGG TCTCAGGGCA ACTCTGTGCA ACTCTGTGGC CCAAAGGGAA TCTACAACTT	sind stop codons)  51   GGGCCCCCGG CAGGGTCGGG CTCCTTCTC GGTGGGAGCA CCTCTTCCACA CCTCTCCACA CTGGCAGCA CCGTGTGGTT TCAGGAGCA GCTGCAGCA CCGTGTGGTT TCAGGAGCAG CCTCACACA CCTCACACAC CCTCACACAC CCTCACACAC CCTCACACAC CCTCACACAC CCTCACACACA	60 120 180 240 300. 360 420 480 540 600 720 780 840 900 960
50 55 60	1 ATGGGAGCC CGCCGACCCC GGCTTCAACT GGATTCTCAG CCCAAGGCTA TGGGTGCCC TGGTTCGGGG AGCTGGCGCA AGCAGGGTT TTAGGTGGAC TTAGGTGGAC ATTGCAGAAT CGCCAGGCCA ATTGCAGAAT CGCCAGGCCA CGCTAGTCAC CGCTAGTCCAC CAGATGCCC CAGATGCCCC CAGATGCCCC	11   GGACGCCAGA CGCTSSTGCC TAGACGCGA TAGACGCGA ATACCAGCCA ACACTACCA CACTGTCCA CACAGATTCG CAGAGAAGGA CCGAATTCT ACTGCCAAGG CAGGAAGCTA CTTATTACCC GTTCCATCTA ATGACACAGA CCATCCTTAA CCTACTTCG	1-3150 (under  21    GTCCCCTCTC GCTGCTGTTG GGCCCAGGA GCCAGGAGTG GTGACCCC CTCAGAGGGA AGCCCATGAG GGAGTATGCA AGGCTTCAGT TTTCTGGCAA CGAGTACCTG TGATGACAGC GGAGTATCTT TGGCTCAGAC CGACTTCGTT	31   CACGCCGTGC CTGCTSSTGC GTACTCTGG ACAGACGGG ACAGACGGG ACTGCAGGTG ATTGAATTTG GAGGGCCTG GACCCGTGG CCTGCAGTTCA GCCGAGTTCA ATCAACCTGG TACCTAGGAT CATCAGATCC ATCAACCTGG CCTGCTGTGC CCTGCTGTGC CCTGCTGCTGC CCTGCTGCAGATCC ATCAACCTAGGAT CACCTAGGAT	41  AGTGCGCTG GGCCCCCGG GCCCCCGG GTCCGCGTTA ACAGCAAAGG TGGATTACA ACAGCATACGT ACAGCATACGT TGGATTACA ACAGCATCGCT ACAGCATCGCT TGCATCTCAC CCAAGACTGG TGTCTGCCAC TTCAGGGGCA ACTCTGTGC CCAAAGGGAA ACTCTGTGC CCAAAGGGAA ACTCTGTGC CCAAAGGAACT TTACAACTT ACGTCAATCG	and stop codons)  51   GGGCCCCGG CAGGGTCGGG CTCCTTCTC GGTGGGAGCA CTCTCGCCT CTCTCGCCT CTCTCGCAG TCCACTGTAC CCTCTCCACA CTGGCAGCA CCGTGTGGTT TCAGGAGCAG GCTGCAGCAT TGTTGGTAA CCTCACTTAC CCTCACTTAC CCTCACTTAC CCTCACTTAC CCTCACTTAC	60 120 180 240 300. 360 420 480 540 600 660 720 780 840 900 960 1020
50 55	1   ATGGGAGCC GCCTTCAACT GGATTCTCAG CCCAAGGCTA CTGGGTGCCA CTGGAGTCCT TGGTTCGGGG AGCTGCCGCA GATAACTTCA GGACAGGGTT TTAGGTGGAC TTAGGTGAAC TCAGGCCA ATTCAGTGGAC ATTCAGTGCAC TTCAGTGCTC GCCCAGGCCA TCAGTGCTC GCCTATGTCA CAGATGCCCT GATGACTTCC	11   GGACGCCAGA CGCTSSTGCC TAGACGCGGA TGGAGTTTTA ATACCAGCA CACTGTCCAG CAGAGAGTTCT ACTGCCAAGG CCGGAATTCT ACTGCCAAGG CAGGAAGCTA ATTACCC CTTCCATCTA ATGACACAGA CCATCCTTAA CCTACTTTAGC TGGTGGGGGC	1-3150 (under  21    GTCCCCTCTC GCCCCAGCA CCGGCCGGGA GCCAGGAGTG GTGCACCCC CTCAGAGGGA AGCCCATGGC GCACTGAGC GGAGTATGCA AGGCTTCAGT TTTCTGGCAA CGAGTACCTG TGATGACAGC AGACTTTGTT TGGCTCAGAG AGACTTTGTT TGGCTCAGAG AGACTTTGTT AGACTCAGT CTATGCAGT ACCCTGCTC	ined sequences of  31   CACGCCGTGC CTGCTSSTGC GTACTCTCGG ACAGACGGGG CTGCAGGTTG ACTCCATCT GACCCGTGG CCCTGCCGCT GCCGGGTTCA GGCCAGATCA GGCAGATCG TACCTAGGAT GCTGGTGTG GCTGGTGTGC ATCAACCTGG ATCTAGATCG ATCGATCGA	41  AGCTGCGCTG GGCCCCCGGG TCAGTGTGCT ACAGCARAGG TGGAGTACAA TGGCATGCGC CGACCTGCTA CAGATTTCAG CCAAGACTGG TCTCAGCGC TCAAGACTGG TCTCACCGC TCAAGGGAA ACTCTGTGGC CCAAAGGGAA TCTACAACTT ACGTCAATGG CCCTGACGG CCCCTGACGG	sind stop codons)  51   GGGGCCCCGG CAGGGTCGGG CTCCTTCTTC GCTGGCTC GTCCTTCCAC CTCTCCACA CCTCTCCACA CCTGGCCAGCA CCGTGTGGTT TCAGGAGCAG CCGTGTGGTT TCAGGAGCAG CCTCACTTAC CTCACTTAC CTCACGGCAAC GCTCACTTAC CTCACGGCAAC CCTCACTTAC CTCACGGCAGA CCTCACTTAC CCTCACGGCAGA GGACGGCCTCAG GCGCCTCAG	60 120 180 240 300 360 420 480 540 660 720 780 840 900 960 1020 1080
50 55 60	1   ATGGGAGCC GGCTGACCCC GGCTTCAACT GGATTCTCAG CCCAAGGCTA TGGGGTGCCA GGTTCGGG AGCTGGCGA GATAACTTCA GGACAGGGTT TTAGGTGGAC ATTGCAGAGCT TCGCCAGGCTA TTCAGTGTGC GGCTATGTCA GGCTAGTGCT GAGGATCGCC CAGATGCCC GAGGTCGCCA GATGACTCC GAGGTCGCCA GAGGTCGCC GAGGTCGCCA GAGGTCGCC GAGGTGGCCA	11   GGACGCCAGA CGCTSSTGCC TAGACGCGA TAGACGCGA ATACCAGCCA ACACTACCA CACTGTCCA CACAGATTCG CAGAGAAGGA CCGAATTCT ACTGCCAAGG CAGGAAGCTA CTTATTACCC GTTCCATCTA ATGACACAGA CCATCCTTAA CCTACTTCG	1-3150 (under  21  i GTCCCCTCTC GCTGCTGTTG GCCCCAGGA CCGGCCGGGA GCCAGGAGTA GTGCACCCC CTCAGAGGGA AGCCCATGAC GGAGTATGCA AGGCTTCAGT TTTCTGGCAA CGAGTACCTG TGATGACAGC AGACTTTGTT TGGCTCAGAC CTATGCAGTA CCCTGCTC CTACCTGCAC CTACCTGCAC CTACCTGCAC	ined sequences of  31    CACGCCGTGC CTGCTSSTGC GTACTCTCGG ACAGACGGGG CTGCAGGGTG TCCTCCATCT GACCCCGTG GCCGAGTTCA GGCAGATCC GCCGAGTTCA GCTGGTGTGC ATCAACCTGG GCTGGTGTGC ATTCGATCCC GCCGCCACAG ATTCGATCCC GCCGCCACAG ATGGATCCG CACCCAGCCA ATGGATCCG CACCCAGCCA CACCCAGCCCA CACCCAGCCCA CACCCAGCCC	41  AGCTGCGCTG GGCCCCCGGG TCAGTGTGCT GTGCTGTCTA TGGCATACAA TGGCATACAC TGGATTCAG CCAAGACTGG TTCAGGGCA TTCAGGGCA ACTCTGTGGC CCAAAGGGAA TCTACAGCTCA TCTACAGCTCA CCAAAGGGAA TCTACAACTT ACGTCAATGG CCCATGACGG CCCCTGACGG GCATAGACCC	and stop codons)  51    GGGCCCCCGG CAGGGTCGGG CTCCTTCTTC GGTGGGAGCA CCTCTTCCAC GTCCTTCAC GTCCTTCAC GTCGTTCAC GTCGTTCAG GTCGTTCAG GTCGTTGCAG TCACTTAC CCTCTCCACA CCGTGTGGTT TCAGGAGCAG CCTCAGACA CCTCAGGCAGA CCTCAGCTAC CTCAGGGGAA GGACGGCTTG GCGGCCTCAG CACGCCCAC CAACGCCCAC	60 120 180 240 300. 360 420 480 540 600 660 720 780 840 900 960 1020
50 55 60	1   ATGGGAGCC CGCCGACCCC GGCTTCAACT GGATTCTCAG CCCAAGGCTA TGGGTGCCC AGCTGCGCA AGCTGGCGCA AGCTGGCGCA ATTGCAGGTT TTAGGTGGAC ATTGCAGAAT CGCCAGGCCA ATTCAGTGGTC ATTCAGTGGTC CGCTAGTCCA CAGATGCCT GATGACTTCA CAGATGCCCT GATGACTTCC GAGGTGGCCA GACCTGGACC GACCTGGACC GACCTGGACC GACCTGGACC GACCTGGACC	11   GGACGCCAGA CGCTSSTGCC TAGACGCGGA TGGAGTTTTA ATACCAGCCA CCCCACACA CACTGTCCAG CACAGATTCG CAGAGAAGGA CCCGAATTCT ACTGCCAAGG CAGGAAGCTA CTTATTACCC GTTCCATCTA ATGACACAGA CCATCCTTAA CCTACTTTAG CGGTTCGGGGGC CGGGTCTACGT TGGCCATGA AGGATGCTA AGGATGCTA	1-3150 (under  21    GTCCCCTCTC GCTGCTGTTG GCCCCAGGA GCCAGGAGTG GTGACCCCC CTCAGAGGGA AGCCCATGAC GCACTGACC GCACTGACC GCACTGACC GCACTGACC GGAGTATCCA TTTCTGGCAA CGACTACCTG TGATGACAC CTATCCAGTC CTACCAGTC CTACCAGTC CTACCTCCAC CTACCTCCAC CAATGATGTG CAATGATGTG CAATGATGTG CAATGATGTG CAATGATGTG CAATGATGTG CAATGATGTG CAATGATGTC	31   CACGCCGTGC CTGCTSSTGC GTACTCTGG GTACTCTGG ACAGACGGG ACAGACGGG ACTGCATCT GACCCGTGG CCCTGCATCT GACCCGTGG CCCTGCAGATCC ATCAACCTGG TACCTAGGAT GCTGGTGTGC ATCAACCTGG ATCAACCTGG ATCGATCC GCCGCCACAG ATGGATCC GCCGCCACAG ATGGATCGG CACCCACCG CGCTTTGGCA CACCCAGCG CACCCAGCG CGCCTTGGGGG GCCATCGGGG GCCATCGGGG	41  AGTGCGCTG GGCCCCCGG GGCCCCCGG GTGCTGTCTA ACAGCAAAGG TGGATTCAG CCAAGACTG GTGCTGCTA ACAGCATCGC TTCAGGGCA TTCAGGGCA ACTCTGCCA TTCAGGGCA ACTCTGCGC CCAAAGGATA ACTCTGCGC CCAAAGGGAA ACTCTGTGC CCAAAGGGAA ACTCTGTGC CCAAAGGGAC CCAAAGGGAC CCCTGACGG GCATTGACG CCCTTTGAC CTCCCTTTGC	and stop codons)  51   GGGCCCCCGG CAGGGTCGGG CTCCTTCTTC GGTGGGAGCA CCTCTGGCTC GTCCTTCCACA CTCTCCACA CTGGCAGCA CCGTGTGGTT TCAGGGAGCA CCGTGTGGTT TCAGGGGAGA CCTCACTTAC CTCACTTAC CTCACTGGGAA GGAGGGCTGAGCCCCCCCCCGGGGTAGACC	60 120 180 240 300 360 420 480 600 660 720 780 840 900 1020 1080 1140 1200 1260
<ul><li>50</li><li>55</li><li>60</li><li>65</li></ul>	1   ATGGGGAGCC GGCTTCAACT GGATTCTCAG CCCAAGGCTA CTGGAGTCCT TGGTTCGGGG AGCTGGCGCA GATAACTTCA GGACAGGGTT TTAGGTGAAC TTAGGTGAAC TTAGGTGAAC TTAGGTGAAC TTAGGTGAC ATTCAGTGGT GGCTATGTCA AGGATGCCC CAGGCCCA CAGGCCCA CAGGCCCA CAGGCCCA CAGGTGGCCC CAGGCCCA CTTACCCTCA CAGCTGGACC CAGCAGGCAC CAGCAGGGAC CAGCAGGGAC	11   GGACGCCAGA CGCTSSTGCC TAGACGCGGA TGGAGTTTTA ATACCAGCCA CACTGTCCAG CAGAGAGTTCT ACTGCCAAGG CAGGAAGCTA ATTATACCC GTTCCATCTA ATGACACGA CCATCTTAC ATGACACGA CCATCTTAC CTGCCATGG CGGTCTACGT CTGCGCGGGC GGGTCTACGT TAGTGCCATGT TAGTGCCATTA TAGTGCCATTA TAGTGCCATCT TAGTGCGCATTA TAGTGTTTTGT	1-3150 (under  21    GTCCCCTCTC GCCCCGGGA CCGGCCGGGA GCCAGGAGT GGCCCATGAG AGCCATGAG AGCCATGAG TTTCTGGCAA AGGCTTCAGT TTTCTGGCAA TGATACAGC GAGATTTGTT TGGTCAGAC CTATGAGAG ACCCTTGCTA ACCCTGCTC CTACCTGCAG ACCCTGCTC CTACCTGCAG ACCCTTGCAG ACCCTTGCTC CTACCTGCAG ACCCTTGCAG ACCCTTGCAG ACCCTTGCAG ACCCTTGCAG ATTTCCTGCAG ATTTCCTGCAG ATTTCCTGCG	ined sequences of  31   CACGCCGTGC CACGCCGTGC CTGCTSSTGC GTACTCTCGG ACAGACGGGG CTGCAGGTG ATTGAATTTG GACCCGTGCCCTT GCCGGTGCAGATCA GGCCAGATCA GGCCAGATCG TACCTAGGAT GCTGGTGTGC ATTCGATCCC GCCGCCACA ATGGATCGA ATGGATCGA CACCAGCCG GCGCCACAG ATGGATCGGA CACCAGCCG GGCTCAGGAG GCCACAGGAG GGCCCAGGAG	41  AGCTGCGCTG GGCCCCCGGG TCAGTGTGCT ACAGCARAGG TGGAGTACAA TGGCATGCGC GGCACCTGCTA CAGATTTCAG CCAAGACTGG TCTCAGGGCA ACTCTGTGGC CCAAAGGGAA ACTCTGTGGC CCAAAGGGAA CCTACAATGG CCAAAGGGAA CCTACAATGG CCCTGACGG GCATAGAGCC CCTCTTTAG CCTCCTTTAGG GGCTGGCTC	sind stop codons)  51   GGGGCCCCGG CAGGGTCGGG CTCCTTCTTC GGTGGGAGCA CCTCTGCCTC GTCCTTCCAG CTCTCCACA CTGGGCAGCA CCGTGTGGTT TCAGGAGCAG CCTCTGCAGA CCTCACTTAC CTCACGGGAAC CCTCACTTAC CTCACGGGAAC CTCACGGGAA CCTCACTTAC CTCACGGGAA CCTCACTTAC CTCACGGGAA CCTCACTTAC CTCACGGGAA CTCACTTAC CTCACGGGAA CTCACTTAC CTCAGGGGAA CTCACTTAC CTCACGGGAAC CACGCCTCAG CACGCCTCAG CACGCCTAG CACGCCTACC CCCCTTGGGG TAGGCCTTCC	60 120 180 240 300 360 420 480 540 660 720 780 840 900 960 1020 1080 1140 1200 1260 1320
50 55 60	1   ATGGGGAGCC GGCTTCAACT GGATTCTCAG CCCAAGGCTA TGGGTGCCA TGGGTGCCA GGATTCTGGGG AGCTGGCGA AGTTGCGGGA ATTGCAGAGTT TTAGGTGGAC TTAGGTGGAC TTCAGTGGTG GGCTATGCAC CAGATGCCCT GAGATGCCCT GAGATGCCCT GAGATGTGCA GACTGGCCA CAGATGCCCT CAGAGGGGAC CTTACCTCA GACCTGGAC CAGCAGGGAC CAGCAGGAC CAGCAGGGAC CAGCAGGGAC CAGCAGGGAC CAGCAGGGAC CAGCAGGGAC CAGCAGGAC CAGCAC CAGCAC CAGCAGGAC CAGCAC CAGCAGGAC CAGCAC CAC	11   GGACGCCAGA CGCTSSTGCC TAGACGCGGA TGGAGTTTTA ATACCAGCCA CACTGTCCAG CAACAGTTCG CAGAGATGCT ACTGCCAAGG CAGGAAGCTA ACTGCCAAGG CAGCAACCTTACA ATGACACGA CCTACTTAG CCTACTTAG CCTACTTAG CCTACTTAG CGGTCTACGT TGGTGGGGGG GGGTCTACGT TAGGTTTTGT AGGCCATGT AGGCCTTTA	1-3150 (under  21    GTCCCCTCTC GCCCCAGCA CCGGCCGGGA GCCAGGAGTA GTGCACCCC CTCAGAGGGA AGCCCATGAC GGAGTATGCA AGGCTTCAGT TTTCTGGCAA AGACTTCAGT TGATGACAGC CTATGACAGC CTATGACAGC AGACTTTGTT TGGCTCAGAC CTATGCAGTA CAATTATGACTAGA TGAGTTTTGGC CAATGATGATGA CATTCCTGGG GGCAGCCAGC	ined sequences of  31    CACGCCGTGC CTGCTSSTGC GTACTCTCGG ACAGACGGGG CTGCAGGTG ATTGAATTTG GAGGAGCCTG GCCAGGTTCA GCCAGGTTCA GCCAGGTTCA GCCAGGTTCA GCCAGGTTCA GCCAGGTTCA GCCAGGTTCA GCCAGGTTCA GCTGGTGTCC GCCGCACAG ATTGATCCC GCCCACAG CCACCCGC GGATTTGGCA CACCCAGCCG CGATTTGGCA CACCCAGCCG CGATTTGGCA CCACCCAGCCG CGATTTGGCA CCACCCAGCCG CGATCCGGGG GCCCACAGCCG CACACCCCAG	41  AGCTGCGCTG GGCCCCCGGG TCAGTGTCTA ACAGCARAGG TGGATACAA TGGCATGCGC CCAGAGATTCAG CCAAGACTG TCTGTCTA ACAGCATGCGC CCAAGACTG TTCAGGGCA ACTCTGTGGC CCAAAGGGAA TCTACAACTT TAGGCATCAATGG CCAAAGGGAA CCTCTTACACCC CCAAAGGGAA TCTACAACTT CCCTTTGGC CCCCTTTGAC CTCCCTTTGG	and stop codons)  51    GGGCCCCCGG CAGGGTCGGG CTCCTTCTTC GCTGGCTC GTCCTTCCAC GTCCTTCCAC GTCGCACAC CCTCTCCACA CCTGGCAGCA CCGTGTGGTT TCAGGAGCAG CCTCACTTAC CTCACGGGAA GCTCACACAC CTCACTTAC CTCAGGGGAA GGACGGCTC CCCCCTGGGG TGGGGAGAC CCCCCTGGGG TGGGGAGAC CCCCCTTCCCCTT	60 120 180 240 300 360 420 480 540 660 720 780 960 1020 1140 1260 1260 1320 1380
<ul><li>50</li><li>55</li><li>60</li><li>65</li></ul>	1   ATGGGAGCC CGCCGACCCC GGCTTCAACT GGATTCTCAG CCCAAGGCTA TGGGGTGCCA ACCTGGAGTCT TAGGTGGAC ATTACTTCA GGACAGGGTT TTAGGTGGAC ATTGCAGAAC ATTGCAGAAC ATTGCAGACT GCCAGGCTC GACTAGCTC GAGTGGCCA CAGAGGGCA CTTACCTCA GACTGGACC CAGCAGGGAC CAGCAGGGAC CAGCAGGAC CAGCAGGCAC CAGCAGGAC CAGCAGGAC CAGCAGGAC CCGAGGAGCC CGAGGAGCC CGAGGAGCC CGAGGAGGCC	11   GGACGCCAGA CGCTSSTGCC TAGACGCGGA TGGAGTTTTA ATACCAGCCA GCCCACACA CACTGTCCAG CAACAGTTCG CAGAGAAGGA CCGAATTTT ATTCTACCAAGG CAGGAAGCTA CTTATTACCC GTTCCATCTA CCATCCTTAA CCATCCTTAA CCATCCTTAA CCTACTTGGG GGGTCTACGT CTGGCCATGA AGGATGGTT AGGATGGTTT AGGATGGTTA AGGATGCTTAA AGGATGCTTAA AGGATGCTTAG AGGATGCTTAG AGGATGCTTAG AGGATGCTTGG GAGACCTGGG GAGACCTGGG	1-3150 (under  21    GTCCCCTCTC GCTGCTGTTG GCGCCAGGAG GCCAGGAGTAG GTGCACCCC CTCAGAGGGA AGCCCATGAGC GGAGTATGCA AGGCTTCAGT TTTCTGGCAA CGAGTACCAG TGATGACAG CTATGAGAG CTATGAGAG AGCCTTCAGA CCCTGCTA ACCCTGCTA ACCCTGCTA TGAGTTTGGC CTACTGCAG TGAGTTTCGT TGAGTTTCGC CAATGATGCAG CTATCCTGCAG TGAGTTTCCTGG ATTTCCTTGG ATTTCCTTGG TGGCAATGGA	ined sequences of  31    CACGCCGTGC CTGCTSSTGC GTACTCTGGG GTACTCTGGG ACTGCAGGTG ATTGAATTTG GAGGGCCTG GCCGAGTTCA GCCGAGTTCA GCCGAGTTCA GCCAGCTG GCCGAGTTCA ATCAACCTGG TACCTAGGAT GCTGGTGTGC GCGCCACAG ATGGATCGA ACCCAGCG GCCACAGGGGGGCCACAG GCCATCGGGG GCCATCGGGG GCCCACGGAG GCCATCGGGG GCCCACGCA CACCCCAG TATCCTGATC TATCCTGATC	41  AGCTGCGCTG GGCCCCCGG GGCCCCCGG GGCCCCCGG GGCCCCCGG GGCCCCGG GGCCCCGGG GGCCCCGGG GGCCCCGGG GGCCCCGGG TGGAGTACAA ACGCAAGACTGG GCACTGCTA CAGATTCAG CCAAGACTGG TCTCAGGGCA ACTCTGTGGC CCAAAGGGA ACTCTGTGGC CCAAAGGGA CCCTGACGG GCATAGAGCC GCTCCTTTGG GGCTGGCTC ACTCCTTTGG GGCTGGGCTC ACTTCTTTGG TGATTGTGGG	and stop codons)  51    GGGCCCCGG CAGGGTCGGG CTCCTTCTC GGTGGGAGCA CTCTTCGCT CTCTCGCTC CTCTCGCAG CTCTCCACA CTGGCAGCA CCGTGTGGTT TCAGGAGCAG CCTCACTTAC CTCACTTAC CTCACTTAC CTCACTTAC CTCACTTAC CTCACTGGAGCAG GCTGCAGCT GTGGGGAA GGACGGCCAC CCCCCTGGGG TGGGGAACC TAAGCCTTCC CTCTCGCCTT GTCCTTTGGT GTCCTTTGGT GTCCTTTGGT GTCCTTTGGT	60 120 180 240 300 360 420 660 660 720 780 840 900 1020 1140 1200 1260 1380 1380 1440
<ul><li>50</li><li>55</li><li>60</li><li>65</li></ul>	1   ATGGGAGCC GCCTACACC GCTTCAACT GGATTCTCAG GCCTAGGCCA CTGGAGTCCT TGGTTCGGGG ACTGGCGCA ACTGGAGTCT TTAGGTGGAC ATTGCAGACT TTAGGTGGAC ATTCAGTGGT GGCTATGTCA GGCTAGTCC GAGTGCCC GAGTGCCC CAGGTGGCCA CTTACCCTCA GACTTGGCC CAGGTGGCC CTTGGACC CAGGTGGCC CTTGGACC CAGGAGGCC CAGGAGGACC CAGGAGAGCC CGAGGAGGCC CGAGGAGACACG CTTGGACACG CGTGGACACG CGTGGACACG CGTGGACACACG CTTGGACACACG CTTGGACACACC CTTGACACC CTTGGACACACC CTTGACACACC CTTGGACACACC CTTGGACACACC CTTGGACACACC CTTGGACACACC CTTGACACACC CTTGGACACACC CTTGGACACACC CTTGGACACC CTTGGACACACC CTTGGACACC CTTGGACACACC CTTGGACACC CTTGGACACC CTTGGACACC CTTGGACACC CTTGCACC CTTCACC	11   GGACGCCAGA CGCTSSTGCC TAGACGCGGA TAGACGCGGA TAGACGCACA CACTGTCCAG CACAGTTCC CAGAGAAGTTCT ACTGCCAAGG CAGGAAGTAC CTTATTACCC GTTCCATCTA ACTGCATCTTA ACTGCATCTTA CCTACTTAC CCTACTTAC CCTACTTAC TAGTGCGGGC GGGTCTACGT TAGTGCATGA AGGATGCTTA TAGTGCTTTCT AGGCATGA AGGATGCTTA TAGTGTTTCT AGCCCTTGT AGCCCCTTGT AGCACCTGGA AGGACCTGGA CAGACCTGGA CTGTGGTATTA	1-3150 (under  21    GTCCCCTCTC GCCCCGCAGCA CCGGCCGGGA GCCAGGAGTG GTGCACCCCC CTCAGAGGGA AGCCCATGGC GCACTGAGC GGAGTATGCA AGGCTTAGT TTTCTGGCAA AGGCTTCAGT TGGCTCAGT TGGCTCAGT CGATGACCTG AGACTTGTT TGGCTCAGC AGACTTTGTT CTACCTGCAG ACCCTGCTC CTACCTGCAG CAATGATTTGG ATTTCCTGGG GGCAGCCAGC CAGGGGCCCCC CAGCCCCCCCCCC	ined sequences of  31    CACGCCGTGC CTGCTSSTGC GTACTCTCGG ACAGACGGG ACTGCAGGTG ATTGAATTTG GAGGACCTGC CCCTGCCGTT GCCGAGTTCA ATCAACCTGG TACCTAGGAT GCTGGTGTGC ATCAACCTGG ATCGATCC ATCGATCCG GCCCACAC ATGGATCGA CACCCAGCGG GCCACCAC GCCACCAC GCCACCAC GCCACCAC GCACTTGGCG GCCACCAC GCCACCAC GCCACCAC GCCACCAC GCCACCAC GCCACCAC GCCACCCAC	41  AGTGCGCTG GGCCCCCGG GCCCCCGGG TCAGTGTGCT ACAGCARAGG TGGATTCAG CGAAGACTGG GCACTGCTA CAGATTTCAG CCAAGACTGG TTCAGGGGCA ACTCTGTGC CCAAAGGGAA ACTCTGTGC CCAAAGGGAA ACTCTGTGC CCAAAGGGAA CCTCTTTGG CCCTTACACTT ACGTCAATTG CCCTTTTGG CCCTTTTGG CCCTTTTGG CCCTTTTTGG CCCTTTTTGG CCCTTTTTGG CCCTTTTTGG CCCTTTTTTGG CCCCTTTTTGG CCCCTTTTTGG CCCCTTTTTGG CCCCTTTTTGG CCCCTTTTTGG CCCCTTTTTGG CCCCTTTTGG CCCCTTTTTGG CCCCTTTTTTGG CCCCTTTTTTTGG CCCCTTTTTTGG CCCCTTTTTTTGG CCCCTTTTTTTGG CCCCTTTTTTGG CCCCTTTTTTGG CCCCTTTTTTGG CCCCTTTTTTGG CCCCTTTTTTTGG CCCCTTTTTTGG CCCCTTTTTTTGG CCCCTTTTTTTGG CCCCTTTTTTGG CCCCTTTTTTGG CCCCTTTTTTGG CCCCTTTTTTGG CCCCTTTTTTGG CCCCTTTTTTGG CCCCTTTTTTTGG CCCCTTTTTTGG CCCCTTTTTTTGG CCCCTTTTTTGG CCCCTTTTTTTGG CCCCTTTTTTTGG CCCCTTTTTTTGG CCCCTTTTTTTGG CCCCTTTTTTTGG CCCCTTTTTTTGG CCCCTTTTTTTGG CCCCTTTTTTTT	and stop codons)  51   GGGCCCCCGG CAGGGTCGGG CTCCTTCTTC GGTGGGAGCA CCTCTGCTC CTCTCGCTC CTCTCCACA CTGGCAGCA CCGTGTGGT TCAGGGAGCA CCGTGTGGT TCAGGGCAA CCTCACTTAC CTCACTTAC CCTCACTTAC CCTCACTTAC CTCACTGGGCAA CCTCACTTAC CTCACTGGGCAA CCTCACTTAC CTCACTGGGCAA CCTCACTTAC CTCACTCAC CTCACTCAC CCTCACTCA	60 120 180 240 300 360 420 480 600 660 720 780 840 900 1020 1080 1140 1260 1320 1320 1340 1500
<ul><li>50</li><li>55</li><li>60</li><li>65</li><li>70</li></ul>	1   ATGGGAGCC GCCTTCAACT GCATTCTCAG CCCAAGGCTA CTGGAGTCCT TGGTTCGGGG AGCTGGCGCA AGATACTTCA GGACAGGGTT TTAGGTGGAC ATTGCAGAAC TTCAGTGGAC ATTGCAGAAC TTCAGTGGTG GCCTATGCAC ATTGCAGAC TTCAGTGGTG GCCTATGCCC CAGGCCCA CATTACCTCA CATTACCTCA CAGTGGGCC CAGGCAGGAC CTGGACC CAGGGAGGAC CAGGGAGGAC CGAGGAGAC CGTGGACAC CGTGGACAC CGTGGACAC CGTGGACAC CGTGGACAC CGTGGACAC CGTGGACAC CGTGGACAC CGTGGACAC CGTGGACACAC ATCTTCCCCC	11   GGACGCCAGA CGCTSSTGCC TAGACGCGGA TGGAGTTTTA ATACCAGCCA CACTGTCCAG CAGCAGATTCT ACTGCCAAGG CAGGAAGCTA ACTGTCCAG CAGGAAGCTA CATCTTACCC GTTCCATCTA ATGACACGA CCATCTTAC CCTGTCCATCTA ATGACACGA CCTACTTAG TGGCCATGG GGGTCTACGT TAGTGCCATGT AGGATGCTTA TAGTGTTTCT AGGACCTGG GAGACCTTGG CATGTTTAA CCTACTGTAAA CCTACTGTAA	1-3150 (under  21    GTCCCCTCTC GCCCCGGGA CCGGCCGGGA GCCAGGAGT GGCCCAGGAGT GGCCCATGAG AGCCATGAG AGCCATGAG TTTCTGGCAA AGGCTTCAGT TTTCTGGCAA AGACTTTGTT TGGTCAGAC CTATGAGAGGT ACCCTGCTC CTACCTGCAG ACCCTGCTC CTACCTGCAG ACCCTGCTC CAATGATTGGG ATTTCTGGG GGCAGCCAGC AGAGGAGGAGGAGGCCCC CCCAGAGGAGG	ined sequences of  31    CACGCCGTGC CTGCTSSTGC GTACTCTCGG ACAGACGGGG CTGCAGGTGGTG ACTGCAGGTG ACTGCAGGTG ACTGCAGGTG GACCCGTGG CCCTGCCGCT GCCGAGTTCA ATCAACCTGG TACCTAGGAT GCTGGTGTGC ATTGATCGATCGATCGACCAG ATGGATCGGA CACCAGGCG GGCCCACAGGGG GGCCCAGGAG TATCCTGAGAT CCCATCGGGG GGCCCACAGGAG CCCATCGGGG CCCACTCGGGG CACACCCCAG CACCCCAGGAG CACACCCCAG CCCATCGTGT CCGAGCTCCA	41  AGCTGCGCTG GGCCCCCGGG TCAGTGTGCT ACAGCARAGG TGGAGTACAA TGGCATGCGC CCAGCATTCAG CCAGATTCAG CCAGATTCAG CCAGATTCAG CCAGACTGG TCAGACTGG TCAGACTGG CCAAAGGGAA ACTCTGTGGC CCAAAGGGAA TCTACAACTG CCCTGACGG GCATACAGC CCCTGACGG GCTCCTTTAG GGCTGGCTC ACTTCTTTGG CCCTTTTGG CCCTTTTTGG CCCTTTTTGG CCGCTAGTGGG	sind stop codons)  51   GGGGCCCCGG CAGGGTCGGG CTCCTTCTTC GGTGGGAGCA CCTCTGCCTC GTCCTTCCAC CTCTCCACA CTGGCCAGCA CCGTGTGGTT TCAGGAGCAG CCTCACTTAC CTCACTTAC CTCACTTAC CTCACTTAC CTCACGGGAA CCTCACTTAC CTCACTCAGGAGCAG GGAGGCCTCAG CACGCCCACC CCCCTGGGG TAAGCCTTCC TTAGCCCTT GTCCTTTGCTTAGC CTCACTTAC CCCCTTGCCTT CCCCTTTGCCTT CTCCCCTTTGCCTT CTCCCTTACC GAACCCTGAC GAACCCTGTG	60 120 180 240 300 360 420 480 540 660 720 780 840 900 960 1020 1080 1140 1200 1320 1380 1440 1500 1500
<ul><li>50</li><li>55</li><li>60</li><li>65</li><li>70</li></ul>	1   ATGGGAGGCC GGCTGACCCC GGCTTCAACT GGATTCTCAG CCCAAGGCTA CTGGAGTCCT TGGTTCGGGG AGCTGGCGCA AGTTGCAGGGTT TTAGGTGGAC ATTGCAGAACT TTAGGTGGAC ATTGCAGAAC TTCAGTGTC GGCTATGTCA GACAGGCTC GACAGGCTC CAGAGCGCC CAGAGCGCC CAGAGGAGCC CAGAGGAGGCC CAGAGGAGGCC CAGGAGAGGCC CAGGAGAAGGCC CTGGACAAGG ATCTTCCCCG GCCTGCATCA	11   GGACGCCAGA CGCTSSTGCC TAGACGCGGA TAGACGCGGA TAGACGCACA CACTGTCCAG CACAGTTCC CAGAGAAGTTCT ACTGCCAAGG CAGGAAGTAC CTTATTACCC GTTCCATCTA ACTGCATCTTA ACTGCATCTTA CCTACTTAC CCTACTTAC CCTACTTAC TAGTGCGGGC GGGTCTACGT TAGTGCATGA AGGATGCTTA TAGTGCTTTCT AGGCATGA AGGATGCTTA TAGTGTTTCT AGCCCTTGT AGCCCCTTGT AGCACCTGGA AGGACCTGGA CAGACCTGGA CTGTGGTATTA	1-3150 (under  21  i GTCCCCTCTC GCCCCAGCA CCGGCCGGGA GCCAGGAGTA GTGCACCCC CTCAGAGGGA AGCCCATGAC GGAGTATGCA AGGCTTCAGT TTTCTGGCAA AGACTTCAGT TGATGACACC CTAACACC CTAACACC CTAACACC AGACTTCAGT TGATGACAC TGATGACAC CTACCTGCAG TGATTTCCAGG GCAGCTAC CAATGATGAT TTCCTGGG GGCAGCCAGC TGGCAATGGA CCCAGGGGCCCAC CCCAGAGGAG CTGCCTCAAT	31   CACGCCGTGC CTGCTSSTGC GTACTCTCGG ACAGACGGGG CTGCAGGTG ATTGAATTTG GAGGAGCCTG CCCTGCCGCT GCCAGGTTCACCTGG ATCAACCTGG ATCAACCTGG ATCAACCTGG CGCGCCACAG ATGGATCCC GCCGCACAG CCCAGCCG CGATTTGGA CCCAGCCG CGATTTGGAT CCCAGCCG CGATTCGGA CCCAGCCG CGATTTGGAT CCCAGCCG CGATTTGGAA CCCAGCCG CGATCGTGG CACACCCAG CCCACAGCG CCCACAGCG CCCACAGCAG CCCACACGTG CCCACAGCAG CCCACCAGC CCCACACGTG CCGAGCTCCAA CCCATCGTGT CCGGAGCTCCAA CCTTCTGGAA	41  AGCTGCGCTG GGCCCCCGGG TCAGTGTCTA ACAGCARAGG TGGATACAA TGGCATGCGC CCAGAGATTCAG CCAAGACTG TCAGAGATTCAG CCAAGACTG TTCAGGGCA ACTCTGTGGC CCAAAGGGAA TCTACAACTT ACGTCAATGG CCATCATTAG CCATCATTGG CCCTTGAC GCTCCTTGAC CTCCTTTGG CTCCTTTGG CCCCTTTGG CCCCTTTGGG CCTCTTTGG CCCTTGTGGC ACTTCTTTGG TGATTGTGGG CCCTAGTGG CCCTTAGAGGGA ACACGTTCC ACACGTTGC ACACGTTGC ACACGTTGC ACACGTTGC ACACGTTGC ACACGGG AACACGTTCC	and stop codons)  51    GGGCCCCCGG CAGGGTCGGG CTCCTTCTTC GGTGGGAGCA CCTCTTGCCT CTCTCGCTC GTCCTTCCAC CTGGCAGCA CCGTGTGGTT TCAGGAGCAG CCGTGTGGTT TCAGGAGCAG CCTCACTTAC CTCACTGGCAAC CCTCACTTAC CTCAGGGGAA GGACGGCTCAG CAGGCCCACC CCCCCTGGGG TGGGGAGAC CTCACTTCC CTCTGCCCTT GTCCTTTGGT CTCCTCACC CTCTCCCTCAC CTCCCTCACC CTCCCCTCGGG GACCCTCGT CGACCCTCGT TGACTCCATT	60 120 180 240 300 360 420 480 600 660 720 780 840 900 1020 1080 1140 1260 1320 1320 1340 1500
<ul><li>50</li><li>55</li><li>60</li><li>65</li></ul>	1   ATGGGAGCC CGCCGACCCC GGCTTCAACT GGATTCTCAG CCCAAGGCTA TGGGGTGCCA CTGGAGTCCT TGGTTCGGG AGCTGGCGA ACTGGAGTTCA CGCAGGGTT TTAGGTGGAC ATTGCAGAGT CGCCAGGCT GAGTGCTC GAGTGCTC GAGTGGCCA CTTACCTCA GACTGGAC CAGCAGGGAC CTTACCTCA GACTGGAC CAGCAGGGAC CTGACCAGCAG CAGGTTCTGC CGAGGAGCAC CGAGGAGCAC CGAGGAGCAC CGGGACACAC GCCTGCATCA GCCTTCACAC GCCTGCATCA GCTTTCACAC	11   GGACGCCAGA CGCTSSTGCC TAGACGCGGA TGGAGTTTTA ATACCAGCCA CACTGTCCAG CAACAGTTCG CAGAGATGCT ACTGCCAAGG CAGGAAGCTA CCTAATTACCA GTTCCACTA ATGACACGA CCTACTTAG CGTCCATCT ATGACACGA CGTCCTTAC ATGACACGA CGGTCTACGT CTGGCCATGA AGGATGCTTA AGGCCCTGTG AGGACTGGT AGCCCTTTG AGCCCTTTACTT ACCTTTGGTATA	1-3150 (under  21    GTCCCCTCTC GCTGCTGTTG GCGCCCGGGA GCCAGGAGTG GTGCACCCC CTCAGAGGGA AGCCCATGAGC GGAGTATGCA AGGCTTCAGT TTTCTGGCAA CGAGTACCTG TGATGACAGC TGATGACAGC TGATGACAGC TGAGTACCTG AGACTTTGTT TGGCTCAGAC CTACCTGCAG TGAGTTTCGTG ACTCTGCAG TGAGTTTCGTG CATTCCTGCAG TGAGTTTCGTG CATGAGTAGAGC CAATGATGT ACTCGCAGC TGGCAATGAG CCCAGAGGGA CCCAGAGGGA CTGCCTCAAT GCTGCAATGA	31    CACGCCGTGC CTGCTSSTGC GTACTCTGGG GTACTCTGGG ACAGACGGG CTGCAGGTG ATTGAATTTG GAGGAGCCTG CCTGCATCT GACCCGTGG CCTGCAGTTCA ATCAACTTG GCCGAGTTCA ATCAACTTG ATCAACTTG ATCAACTTG ATCAACTGGTGTGC GCCGCCACAG ATGGATCCC GCCGCCACAG ATGGATCGG ATCGATCGG GCCCACGGGG GCCCACGGGG GCCCACGGGG GCCCACGGGG CACACCCCAG CCATCGGGG GCCCACGGAG CACACCCCAG CCATCGTGT CCCATCGTGT CCCATCGTGT CCGATCTGAA CAGAAGCAGA	41  AGCTGCGCTG GGCCCCGGG GGCCCCGGG GGCCCCGGG GGCCCCGGG TGGGTGTCTA ACAGCAAAGG TGGATTCAG CCAAGACTGC GCACTGCTA CAGATTCAG CCAAGACTGC TTCAGGGCA ACTCTGTGC CCAAAGGCA ACTCTGTGC CCAAAGGCA ACTCTTTGG GCATAGACT ACGTCAATGG CCCTTAGAC CTCCTTTGG GGCTGCTT TCTTTTGG TGATTCTTTGG TGATTGTGGC CCGCTAGAGG CCGCTAGAGG ACACGTTCC AGGGAGGCG ACACGTTCC AGGGAGGCG AACACGTTCC AGGGAGGGGT	and stop codons)  51    GGGCCCCGG CAGGGTCGGG CTCCTTCTC GGTGGGAGCA CCTCTGTCT CTCTCGCTC CTCTCGCTC CTCTCCACA CTGGCAGCA CCGTGTGGTT TCAGGAGCAG CCTCACTTAC CCTCACTTAC CCTCACTTAC CCTCACTTAC CCTCACTTAC CCTCACTTAC CTCAGGGAA GGACGGGCTCAG CAGGCCCAC CCCCTGGGG TGGGGAACC TAAGCCTTCC CTCTCCCTT GTCCTTTGGT CTCCCTTAGT CTCCCTCACT GAACCCTCAC GAACCCTAGT TGCCTTAGT TGCCTCACT TGCCTTAGT TGCCTTAGT TGCCTCACT GAACCCTAGT TGCCTCACT TGCCCTCACT TGCCCTCACC TGCCCCGGCCA	60 120 180 240 300 360 420 480 540 660 720 780 960 1020 1140 1200 1260 1320 1380 1440 1500 1560 1620
<ul><li>50</li><li>55</li><li>60</li><li>65</li><li>70</li></ul>	Coding sequence  1	11   GGACGCCAGA CGCTSSTGCC TAGACGCGGA TAGACGCGGA ATACCAGCCA CACTGTCCAG CACAGTTCC CAGAGAAGTTCT ACTGCCAAGG CAGGAAGTAC CTTATTACCC GTTCCATCTA ATGACACAG CCTACTTAC CCTACTTAC CCTACTTAC CTACTTTAC TAGTCCATGA AGGATGGCT AGGCCATGA AGGATGCTTA TAGTGCTACGT CTGGCATGA AGGATGCTTA CCTACTTAG CCTCTTAGA CCTTTAGATA CCTTTAGATA CCTCTTAGATA CCTCTAGGCA	1-3150 (under  21    GTCCCCTCTC GCCCCGCAGCA CCGGCCGGGA GCCAGGAGTG GTGCACCCCC CTCAGAGGGA AGCCCATGGC GGAGTATGCA AGGCTACCA TTTCTGGCAA AGGCTTCAGT TGGTCAGAC TGATGACAGC CTATGCAGT GATGACAGC AGACTTGTT TGGCTCAGAC ACCCTGCTC CTACCTGCAG GGCAGCCAGC CAATGATGTG ATTTCCTGGG GGCAGCCAGC CAGGGGCCGC CCAGGGGAG CTGCCTCAAT GCTGCATCGA GCTGCATCG GGCAACCCTG GCGCACCTG GCGCACCTG GCGCACCTG GCGCACCTG GCGCACCTG GCGCACCTG GCGCACCTG GCGCACCTG GCGCACCTG GCGCACCCTG	31   CACGCCGTGC CTGCTSSTGC GTACTCTGG ACAGACGGG ACAGACGGG ATTGAATTTG GAGGACCTGC CCTTCCATCT GCCGAGTTCA ATCAACTTGG ACCAGACTCG ACCAGACTCG ATCAACCTGG ATCAACCTGG ATCGATCC GCCGCCACAC ATGGATCGA ATGGATCGA ATGCACCAGCG GCCACCAGGAG CACACCCAGGAG CACCACGGGG GCCCACAGAG CACCAGACCCAGAC CCCATCTGTT CGGAGCTGCA CCCATCTGTT CGGAGCTGCA CCCATCACCC ACCAGACCCAAC CCAGAGCAAC ACCCAGAACCCAA	41  AGTGCGCTG GGCCCCCGG GCCCCCGGG TCAGTGTGCT ACAGCARAGG TGGATTCAG CGAAGACTG GGACTGCGA CGAGATTCAG CCAAGACTG TTCAGGGCA ACTCTGTGC CCAAAGGGAA ACTCTGTGC CCATCACACT ACGCTACAG CCCTGACGG GCATAGACCT GCTCCTTTGA CCTCCTTTGAC CCTCCTTTTGG CCCTTGAC CCTCCTTTTGG CCCTTTTGG CCCTTTTTGG CCCCTTTTTGG CCCTTTTTGG CCCCTTTTGG CCCCTTTTGG CCCTTTTTGG CCCCTTTTGG CCCCTTTGGG CCCCTTTGGG CCCCTTTGGG CCCCTTTGGG CCCCTTTGGG CCCCTTTGGG CCCCTTTGGG CCCCTTTGGC CCCCTTTGGG CCCCTTTGGC CCCCTTTTGGC CCCCTTTTGGC CCCCTTTTGGC CCCCTTTTGGC CCCCTTTTTGGC CCCCTTTTTGGC CCCCTTTTTGGC CCCCTTTTTGGC CCCCTTTTTGGC CCCCTTTTTGGC CCCCTTTTTTGGC CCCCTTTTTTGGC CCCCTTTTTTGGC CCCCTTTTTTGGC CCCCTTTTTTGGC CCCCTTTTTTGGC CCCCTTTTTTGGC CCCCTTTTTTTGGC CCCCTTTTTTTT	and stop codons)  51    GGGCCCCGG CAGGGTCGGG CTCCTTCTC GGTGGGAGCA CCTCTGTCT CTCTCGCTC CTCTCGCTC CTCTCCACA CTGGCAGCA CCGTGTGGTT TCAGGAGCAG CCTCACTTAC CCTCACTTAC CCTCACTTAC CCTCACTTAC CCTCACTTAC CCTCACTTAC CTCAGGGAA GGACGGGCTCAG CAGGCCCAC CCCCTGGGG TGGGGAACC TAAGCCTTCC CTCTCCCTT GTCCTTTGGT CTCCCTTAGT CTCCCTCACT GAACCCTCAC GAACCCTAGT TGCCTTAGT TGCCTCACT TGCCTTAGT TGCCTTAGT TGCCTCACT GAACCCTAGT TGCCTCACT TGCCCTCACT TGCCCTCACC TGCCCCGGCCA	60 120 180 240 300 360 420 480 600 660 720 840 900 1020 11200 1260 1320 1320 1380 1440 1500 1560 1680 1740
<ul><li>50</li><li>55</li><li>60</li><li>65</li><li>70</li></ul>	1   ATGGGAGGCC GGCTGACCCC GGCTTCAACT GGATTCTCAG CCCAAGGCTA CTGGAGTCCT TGGTTCGGG AGCTGGCGA GATACTTCA GGACAGGGTT TTAGGTGGAC TTAGGTGAC TTAGGTGGAC TTAGGTGGTC GGCTATGTCA GACAGGGTT CAGAGTCGCC CAGAGGCA CTTACCTCA GACTGGAC CAGAGGAG CAGGTTCTCC CAGAGGAGGC CTGCACCAGGCAC CTGCACCAGGCAC CTTCCCCCG GCTTCCCCG GCTTCCCCG CCTGCATCA GCTTCCCCG CCTGCATCA CCTGTCCCGA	11   GGACGCCAGA CGCTSSTGCC TAGACGCGA TGGAGTTTTA ATACCAGCCA CACTGTCCAG CAACAGTTCG CAGAGAGAGGA ACGCTAATCT ACTGCCAAGG CAGGAGACTAC CATCTTATTACCACTAC CATCCTTAA CCATCCTTAA CCATCCTTAA CCATCCTTAA TGGCCATGA AGGATGCTT AGGCCATGA AGGATGCTT AGGCCTTTG GAGACCTGG CTGTGGTATAT ACCTTGGTATAT ACCTTGGTATAT AGCCCTTTG GAGACCTGG CCATGGTATAT CCATCTTCAC CCTCCAGCA CCTTCCAGCA CCTCCAGCA CCAGGAGAGAT TTCACATCGC	1-3150 (under  21  i GTCCCCTCTC GCCCCAGGA CCGGCCGGGA GCCAGGAGTA GTGCACCCC CTCAGAGGGA AGCCCATGAC GGAGTATGCA AGGCTTCAGT TTTCTGGCAA AGGCTTCAGT TATATGACAGC CTAACAGC CTAACAGC CTAACAGC CTAACAGC CTACTGCAG AGACTTTGT TGGCTCAGA CCCTGCTC CTACCTGCAG CAACATGAGA CCGGAGCCAGC TGGCAACCCTG CCAGGACCCAG GGCAACCCTG GGAGATCTAC TCTCAACTTC TCTCAACTTC	ined sequences of  31    CACGCCGTGC CTGCTSSTGC GTACTCTGG GTACTCTGG ACAGGGGG CTGCAGGTG ATTGAATTTG GAGGAGCCTG GACCCCGTGG CCTGCCGTTG GCCGAGTTCA ATCAACCTGG TACCTAGGAT GCTGGTGTGC ATTGGATCCC GCGCCACAG ATGGATCGA ATGGATCGA GCCACAGCGG GGCCACAG GCCACAGGGG GGCCAGAG TATCCTGGTGT CCCATCGTGT CCCATCGTGT CCGAGCTGCA GCTTCTGGAA CCCAGACCC CTCAGGAACG TCCTTGGAACG TCCTTTGGAACG TCCTTGGAACG TCCTTTGGAACG TCCTTTTGGAACG TCCTTTTTGAACG TCCTTTTTTTTTT	41  AGTGCGCTG GGCCCCGGG GGCCCCGGG TCAGTGTGTTA ACAGCAAAGG TGGAGTACAA ACAGCATCAG CCAAGACTGC TCTCTGCCAC TCTCAGGGCA ACTCTGTGCC CCAAAGGGA TCTACAACTT ACGTCAATGG CCCTGACG GCTTCTTGG GGCTAGGGC TCCTTTGG GGCTAGGGC TCTTTTGG TGATTGTGGG CCGCTAGGGG AACACGTTCC TGCTTAGAGGC TGCTTAGAGGG AACACGTTCC AGGGAGGGG AACACGTTCC AGGGAGGGCT TGCTCATCAC AGTCCAACCCC CCCAAGCCCC	and stop codons)  51  GGGCCCCGG CAGGGTCGGG CTCCTTCTTC CTCTGGGAGCA CCTCTTCCACA CTCTCCACA CCTCTCCACA CCGGGAGCA CCTCTCCACA CCGGGAGCA CCTCTCCACA CCGGGCAGCA CCTCACCTTAC CTCACGGGAGCA CCTCACCTTAC CTCACGGGAA GGACGGCTCAG CCCCCTGGGG TGGGGAGAC CCCCTTGGGT TGACCCTTC CTCTCCCTT CTCCTTAC CTCCCTTAC CAACCCTGT TGACTCCATT ACGGCGGCA GAATGGGGCA GAATGGGGCA CAATGGGCAAA AGTGGACAAA AGTGGACAAC	60 120 180 240 300 360 420 540 600 660 720 780 840 900 1020 1140 1200 1260 1320 1440 1500 1500 1620 1680 1740 1800 1860
<ul><li>50</li><li>55</li><li>60</li><li>65</li><li>70</li></ul>	Coding sequence  1	11   GGACGCCAGA CGCTSSTGCC TAGACGCGGA TAGACGCGGA TAGACACTTCAG CACTGTCCAG CACAGAAGGA CCCGAATTCT ACTGCCAAGG CAGAGAAGGA CCTATTACCC GTTCCATCTA ATGACACAG CCTACTTAG CCTACTTAG CCTACTTAG CGGCCACTA AGGATCGCTA AGGATCTTAG CGTTCGTATA CCTACTTTAG CGTTCGTATT TAGTGTTTCT TAGTGTTTCT TAGTGTTTCA CCATGTTCAT CCATGTTCAT CCATGTTCAT CCATGTTCAC CCTCCAGGCA CCTCCAGGCA CCTCCAGGCA CCTCCAGGCA CCAGCCCCT TTCCAACCCC GCCAGCCCCCT	1-3150 (under  21    GTCCCCTCTC GCGCCGGGA GCCAGCACA CCGGCCGGGA AGCCCATGGC CTCAGAGGGA AGCCCATGGC GGAGTATGGA TTTCTGGCA AGGCTTAGT TTTCTGGCA AGACTTTGTT TGGTCAGAC CTACCTGCAC CTACCTGCAC CAATGATGT ACCCTGCTC CAATGATGTA ATTCCTGGG GGCAGCCAGC CCAGAGGAC CTGCCAGAGGAC CTGCCAACTGCAC CTGCTCAACTTG GAGACTTTAG ATTCCTGGG GCCAGACCAC CTGCACCAC CCAGAGGACCAC CCAGAGGACCAC CTGCACTAGAC CTGCCAACTAG CTGCAACTTC GAAGATCTAC ACATTATCAG	31    CACGCCGTGC CTGCTSSTGC GTACTCCGG GTACTCCGG ACAGACGGGG ACAGACGGGG ACTGCAGGTG ATTGAATTTG GAGGGCCTG CCCTGCCGT GCCGGGTTCA ATCAACCTGG ATCAACCTGG ATCAACCTGG ATCAACCTGG ATCGATCTG GCCGCCACAG ACCCCAGG GCCACAGGGG CACACCCCAG GCCATCGGGG CACACCCCAG GCCATCGGGG CACACCCCAG CACCCCAG CCCATCGTGT CCCATCGTGT CCCATCGTGT CCCATCGTGT CCCATCGTGT CCCATCGTGT CCCATCGTGT CCGAGACCC CTCAGGAACG CTCAGGAACG ACCCAGACCC CTCAGGAACG ACCCAGACCC CTCAGGAACG ACCCAGACCC AGCAACACCC AGCAACACC AGCAACACCC AGCAACACC AGCAACACCC AGCAACACC AGCAACACCC AGCAACACC AGCAACAC AGCAACACC AGCAACACC AGCAACACC AGCAACAC AGCAACACC AGCAACAC A	41  AGCTGCGCTG GGCCCCCGG GCCCCCGGG TCAGTGTGCTA ACAGCAAAGG TGGATTCAG CCAAGACTG CCAAGACTG CCAAGACTG TTCAGGGCA ACTCTGCCA TCAGGGCA ACTCTGTGC CCAAAGGGAA ACTCTGTGC CCCTAACG GCTCCTTTGG GCTTCTTTGG GCTTCTTTGG TGCTTCTTTGG TGCTTCTTTGG TGATTGTGGC CCGCTAGTGC ACTCTTTGG TGATTGTGGG CCGCTAGTGC ACTCTTTGG TGATTGTGGG CCGCTAGTGC ACTCTTTGG TGATTGTGGG CCGCTAGTGC ACTCTTTGG TGATTGTGGG CCGCTAGTGC ACTCTTTGG ACACGTTCC AGGAAGACGCT TCCCAAGCCCC GGATAGAGGA	and stop codons)  51   GGGGCCCCGG CAGGGTCGGG CTCCTTCTTC GGTGGGAGCA CCTCTGCCTC GTCCTTCCAC GTCGTCGCTC GTCCTTCCAC CTGGCCACA CCGTGTGGTT TCAGGAGCAG CCTCACTTAC CTCACTTAC CTCACTTAC CTCACGGGAA CCTCACTTAC CTCACGGGAA CCTCACTTAC CTCACTCAC CCCCCTGGGG TGGGGAGAC TAGCCCTCC CTCTGCCCTT GTCCTTTGGT CTCCCTCAC GAACCCTTCC GAACCCTTCC CTCTGCCCTT GCCCTCAC GAACCCTGTG TGACTCCATT ACGCCGGCA CACGCCGGCAA CCTCACTTAC CTCACC GAACCCTGTG TGACTCCATT ACGCCGGCAAA	60 120 180 240 300 360 420 480 660 720 780 840 900 1020 11200 1200 1200 1200 1200 1320 132

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                                                                               2100
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                                                                               2160
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                                                                               2640
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                                                                               2820
15
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                                                                               3000
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                                                                              3060
        TACAAGCTTG GATTCTTCAA ACGCTCCCTC CCATATGGCA CCGCCATGGA AAAAGCTCAG
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                                                                                 120
30
         LESSLSSEG EEPVEYKSLQ WFGATVRAHG SSILACAPLY SWRTEKEPLS DPVGTCYLST
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                                                                                 300
         GYVTILNGSD IRSLYNFSGE QMASYFGYAV AATDVNGDGL DDLLVGAPLL MDRTPDGRPQ
                                                                                 360
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         QQGVVFVFPG GPGGLGSKPS QVLQPLWAAS HTPDFFGSAL RGGRDLDGNG YPDLIVGSFG
                                                                                 480
         VDKAVVYRGR PIVSASASLT IFPAMFNPEE RSCSLEGNPV ACINLSFCLN ASGKHVADSI
                                                                                 540
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LSPIHIALNF SLDPQAPVDS HGLRPALHYQ SKSRIEDKAQ ILLDCGEDNI CVPDLQLEVF
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                                                                                 660
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         SFRLSVEAQA QVTLNGVSKP EAVLFPVSDW HPRDQPQKEE DLGPAVHHVY ELINQGPSSI
         SQGVLELSCP QALEGQQLLY VTRVTGLNCT TNHPINPKGL ELDPEGSLHH QQKREAPSRS
                                                                                 900
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                                                                                 960
        YKALKMPYRI LPRQLPQKER QVATAVQWTK AEGSYGVPLW IIILAILFGL LLLGLLIYIL 1020
45
        YKLGFFKRSL PYGTAMEKAQ LKPPATSDA
                                                   SEQ ID NO:293 LBH4 DNA SEQUENCE
        Nucleic Acid Accession #: BC001291
50
                          44-541 (start and stop codons are underlined)
        Coding sequence:
            11
                  21
                        31 41
                                    51
55
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       AGATCCAGAG GACTCCCAGC GAACGGACGA GGGTGACAAT AGAGTGTGGT GTCATGTTTG 180
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       CTGCGTTATA GCGGCCGTGA AAATATTTCC ACGTTTTTTC ATGGTTGCGA AGCAGTGCTC 300
60
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GGGATGGGAG AGTGGGGATC AGGTGCAGTT GGCTCTTAAC CCTCAAGGGT TCTTTAACTC 720
ACATTCAGAG GAAGTCCAGA TCTCCTGAGT AGTGATTTTG GTGACAAGTT TTTCTCTTTG 780
       AAATCAAACC TTGTAACTCA TITATTGCTG ATGGCCACTC TTTTCCTTGA CTCCCCTCTG 840
       CCTCTGAGGG CTTCAGTATT GATGGGGAGG GAGGCCTAAG TACCACTCAT GGAGAGTATG 900
TGCTGAGATG CTTCCGACCT TTCAGGTGAC GCAGGAACAC TGGGGGAGTC TGAATGATTG 960
70
       GGGTGAAGAC ATCCCTGGAG TGAAGGACTC CTCAGCATGG GGGCAGTGG GGCACACGTT 1020
       AGGGCTGCCC CCATTCCAGT GGTGGAGGCG CTGTGGATGG CTGCTTTTCC TCAACCTTTC 1080
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       ACCAGCTGGC ACAGGTGCAC AGATTCATAA ATTCCCACAC GTGTGTGTTC AACATCTGAA 1200
75
       ACTTAGGCCA AGTAGAGAGC ATCAGGGTAA ATGGCGTTCA TTTCTCTGTT AAGATGCAGC 1260
       CATCCATGGG GAGCTGAGAA ATCAGACTCA AAGTTCCACC AAAAACAAAT ACAAGGGGAC 1320
       ΤΤΟΑΛΑΛΟΤΤ CACGAAAAAA ΑΛΑΛΑΑΑΑΑΑ ΑΛΑΛΑΑΑΑΑ ΑΛΑΛΑΑΑΑΑ ΑΛΑ
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#### SEQ ID NO:294 LBH4 Protein sequence; Protein Accession #: AAH01291

5 1 11 21 31 41 51 1 1 MALLALLUV ALPRVWTDAN LTARQRDPED SQRTDEGDNR VWCHVCEREN TFECONPRRC 60 KWTEPYCVIA AVKIFPRFFM VAKQCSAGCA AMERPKPEEK RFILEEPMPF FYLKCCKIRY 120 CNLEGPPINS SVFKEYAGSM GESCGGLWLA ILLLLASIAA GLSLS

15

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It is understood that the examples described above in no way serve to limit the true scope of this invention, but rather are presented for illustrative purposes. All publications, sequences of accession numbers, and patent applications cited in this specification are herein incorporated by reference as if each individual publication or patent application were specifically and individually indicated to be incorporated by reference.

## WHAT IS CLAIMED IS:

A method of detecting a prostate cancer-associated transcript in a cell 1 from a patient, the method comprising contacting a biological sample from the patient with a 2 polynucleotide that selectively hybridizes to a sequence at least 80% identical to a sequence 3 4 as shown in Tables 1-16. 2. The method of claim 1, wherein the polynucleotide selectively 1 hybridizes to a sequence at least 95% identical to a sequence as shown in Tables 1-16. 2 1 3. The method of claim 1, wherein the biological sample is a tissue 2 sample. 4. The method of claim 1, wherein the biological sample comprises 1 isolated nucleic acids. 2 1 5. The method of claim 4, wherein the nucleic acids are mRNA. 6. The method of claim 4, further comprising the step of amplifying 1 2 nucleic acids before the step of contacting the biological sample with the polynucleotide. 7. The method of claim 1, wherein the polynucleotide comprises a 1 2 sequence as shown in Tables 1-16. The method of claim 1, wherein the polynucleotide is labeled. 1 8. The method of claim 8, wherein the label is a fluorescent label. 9. 1 The method of claim 1, wherein the polynucleotide is immobilized on 10. 1 2 a solid surface. The method of claim 1, wherein the patient is undergoing a therapeutic 1 11. 2 regimen to treat prostate cancer. 1 12. The method of claim 1, wherein the patient is suspected of having 2 prostate cancer.

1 13. A method of monitoring the efficacy of a therapeutic treatment of 2 prostate cancer, the method comprising the steps of: 3 (i) providing a biological sample from a patient undergoing the therapeutic 4 treatment; and 5 (ii) determining the level of a prostate cancer-associated transcript in the biological sample by contacting the biological sample with a polynucleotide that selectively 6 hybridizes to a sequence at least 80% identical to a sequence as shown in Tables 1-16, 7 8 thereby monitoring the efficacy of the therapy. 14. The method of claim 13, further comprising the step of: (iii) comparing 1 2 the level of the prostate cancer-associated transcript to a level of the prostate cancer-3 associated transcript in a biological sample from the patient prior to, or earlier in, the 4 therapeutic treatment. 1 15. The method of claim 13, wherein the patient is a human. 16. A method of monitoring the efficacy of a therapeutic treatment of 1 2 prostate cancer, the method comprising the steps of: 3 (i) providing a biological sample from a patient undergoing the therapeutic 4 treatment; and 5 (ii) determining the level of a prostate cancer-associated antibody in the biological sample by contacting the biological sample with a polypeptide encoded by a 6 polynucleotide that selectively hybridizes to a sequence at least 80% identical to a sequence 7 8 as shown in Tables 1-16, wherein the polypeptide specifically binds to the prostate cancer-9 associated antibody, thereby monitoring the efficacy of the therapy. 1 **17**. The method of claim 16, further comprising the step of: (iii) comparing 2 the level of the prostate cancer-associated antibody to a level of the prostate cancer-3 associated antibody in a biological sample from the patient prior to, or earlier in, the 4 therapeutic treatment. 1 18. The method of claim 16, wherein the patient is a human.

1		19.	A method of monitoring the efficacy of a therapeutic treatment of
2	prostate cancer	r, the m	ethod comprising the steps of:
3	•	(i) prov	viding a biological sample from a patient undergoing the therapeutic
4	treatment; and		
5		(ii) det	ermining the level of a prostate cancer-associated polypeptide in the
6	biological sam	ple by o	contacting the biological sample with an antibody, wherein the antibody
7	specifically bin	nds to a	polypeptide encoded by a polynucleotide that selectively hybridizes to
8	a sequence at l	east 80°	% identical to a sequence as shown in Tables 1-16, thereby monitoring
9	the efficacy of	the the	гару.
1		20.	The method of claim 19, further comprising the step of: (iii) comparing
2	the level of the		te cancer-associated polypeptide to a level of the prostate cancer-
3		-	e in a biological sample from the patient prior to, or earlier in, the
			e in a biological sample from the patient prior to, or carner in, the
4	therapeutic tre	аппсис.	
1		21.	The method of claim 19, wherein the patient is a human.
1		22.	An isolated nucleic acid molecule consisting of a polynucleotide
2	sequence as sh	own in	Tables 1-16.
1		23.	The nucleic acid molecule of claim 22, which is labeled.
1		24.	The nucleic acid of claim 23, wherein the label is a fluorescent label
1		25.	An expression vector comprising the nucleic acid of claim 22.
1		26.	A host cell comprising the expression vector of claim 25.
1		27.	An isolated polypeptide which is encoded by a nucleic acid molecule
2	having polynu	cleotide	e sequence as shown in Tables 1-16.
1		28.	An antibody that specifically binds a polypeptide of claim 27.
1		29.	The antibody of claim 28, further conjugated to an effector component.

1	50.	The antibody of claim 29, wherein the effector component is a
2	fluorescent label.	
1	31.	The antibody of claim 29, wherein the effector component is a
2	radioisotope or a cy	rtotoxic chemical.
1	32.	The antibody of claim 29, which is an antibody fragment.
1	33.	The antibody of claim 29, which is a humanized antibody
1	34.	A method of detecting a prostate cancer cell in a biological sample
2	from a patient, the 1	method comprising contacting the biological sample with an antibody of
3	claim 28.	
1	35.	The method of claim 34, wherein the antibody is further conjugated to
2	an effector compone	ent.
1	36.	The method of claim 35, wherein the effector component is a
2	fluorescent label.	
1	37.	A method of detecting antibodies specific to prostate cancer in a
2		comprising contacting a biological sample from the patient with a
3		d by a nucleic acid comprises a sequence from Tables 1-16.
1	38.	A method for identifying a compound that modulates a prostate cancer-
2		ide, the method comprising the steps of:
3		ontacting the compound with a prostate cancer-associated polypeptide, the
4	polypeptide encode	d by a polynucleotide that selectively hybridizes to a sequence at least
5	80% identical to a se	equence as shown in Tables 1-16; and
6	(ii) d	etermining the functional effect of the compound upon the polypeptide.
1	39.	The method of claim 38, wherein the functional effect is a physical
2	effect.	

The method of claim 38, wherein the functional effect is a chemical

1

2

effect.

40.

1	41.	The method of claim 38, wherein the polypeptide is expressed in a
2	eukaryotic host cell	or cell membrane.
1	42.	The method of claim 38, wherein the functional effect is determined by
2	measuring ligand bi	nding to the polypeptide.
1	43.	The method of claim 38, wherein the polypeptide is recombinant.
1	44.	A method of inhibiting proliferation of a prostate cancer-associated
2	cell to treat prostate	cancer in a patient, the method comprising the step of administering to
3	the subject a therape	eutically effective amount of a compound identified using the method of
4	claim 38.	
1	45.	The method of claim 44, wherein the compound is an antibody.
1	46.	The method of claim 45, wherein the patient is a human.
1	47.	A drug screening assay comprising the steps of
2	(i) ad	ministering a test compound to a mammal having prostate cancer or a
3	cell isolated therefro	om;
4	(ii) co	omparing the level of gene expression of a polynucleotide that selectively
5	hybridizes to a seque	ence at least 80% identical to a sequence as shown in Tables 1-16 in a
6	treated cell or mamr	nal with the level of gene expression of the polynucleotide in a control
7	cell or mammal, who	erein a test compound that modulates the level of expression of the
8	polynucleotide is a c	andidate for the treatment of prostate cancer.
1	48.	The assay of claim 47, wherein the control is a mammal with prostate
2	cancer or a cell there	efrom that has not been treated with the test compound.
1	49.	The assay of claim 47, wherein the control is a normal cell or mammal

1 50. A method for treating a mammal having prostate cancer comprising 2 administering a compound identified by the assay of claim 47. 1 51. A pharmaceutical composition for treating a mammal having prostate 2 cancer, the composition comprising a compound identified by the assay of claim 47 and a 3 physiologically acceptable excipient. 1 52. The method according to claim 1, wherein said biological sample is 2 contacted with a plurality of polynucleotides comprising a first polynucleotide that 3 selectively hybridizes to a sequence at least 80% identical to a first sequence as shown in 4 Tables 1-16; and a second polynucleotide that selectively hybridizes to a second sequence at 5 least 80% identical to a second sequence as shown in Tables 1-16. 53. A method according to claim 52, wherein the plurality of 1 2 polynucleotides comprises a third polynucleotide that selectively hybridizes to a sequence at 3 least 80% identical to a third sequence as shown in Tables 1-16.. 1 54. A method of detecting a prostate cancer associated transcript, the 2 method comprising contacting a biological sample from the patient with a plurality of polynucleotides wherein at least two of said polynucleotides selectively hybridize to a 3 4 difference sequence at least 80% identical to a sequence as shown in Tables 1-16. 55. 1 A method of detecting a prostate cancer, the method comprising the 2 steps of: 3 (i) providing a biological sample from a patient; (ii) contacting the biological sample with a first polynucleotide that selectively 4

- hybridizes to a sequence at least 80% identical to a first sequence as shown in Tables 1-16 to determine the level of a prostate cancer-associated transcript in the biological sample; and with a second polynucleotide that selectively hybridizes to a second sequence at least 80% identical to a sequence not shown in Tables 1-16; wherein the expression of said second
- 9 sequence is not substantially changed in prostate cancer, to determine the level of expression
- of a control transcript in the biological sample;

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11 (iii) comparing the level of the prostate cancer-associated transcript to a level 12 of the normal tissue associated transcript in the biological sample. 1 56. A method of quantitating a prostate cancer-associated transcript in a 2 cell from a patient, the method comprising contacting a biological sample from the patient 3 with a polynucleotide that selectively hybridizes to a sequence at least 80% identical to a 4 sequence as shown in Tables 1-16. 57. The method of claim 56, wherein the polynucleotide selectively 1 2 hybridizes to a sequence at least 95% identical to a sequence as shown in Tables 1-16. 1 58. The method of claim 56, wherein the biological sample is a tissue 2 sample. 59. 1 The method of claim 56, wherein the biological sample comprises 2 isolated nucleic acids. 60. The method of claim 56, wherein the nucleic acids are mRNA. 1 61. 1 The method of claim 59, further comprising the step of amplifying 2 nucleic acids before the step of contacting the biological sample with the polynucleotide. 1 62. The method of claim 56, wherein the polynucleotide comprises a sequence as shown in Tables 1-16. 2 1 63. The method of claim 56, wherein the polynucleotide is labeled. 1 64. The method of claim 63, wherein the label is a fluorescent label. 1 65. The method of claim 56, wherein the polynucleotide is immobilized on a solid surface. 2 . 66. The method of claim 56, wherein the patient is undergoing a 1 2 therapeutic regimen to treat metastatic prostate cancer. 1 67. The method of claim 56, wherein the patient is suspected of having

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metastatic prostate cancer.

1	68. A biochip comprising a plurality of polynucleotides that selectively
2	hybridize to a sequence at least 80% identical to a sequence as shown in Tables 1-16.
1	69. A method of screening drug candidates comprising:
2	i) providing a cell that expresses an expression profile gene selected from the
3	group consisting of an expression profile gene set forth in Tables 1-16 or fragment thereof;
4	ii) adding a drug candidate to said cell; and
5	iii) determining the effect of said drug candidate on the expression of said
6	expression profile gene.
1	70. A method according to claim 59 wherein said determining comprises
2	comparing the level of expression in the absence of said drug candidate to the level of
3	expression in the presence of said drug candidate.
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